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<p>(21) International Application Number: PCT/GB99/02444</p> <p>(22) International Filing Date: 27 July 1999 (27.07.99)</p> <p>(30) Priority Data:</p> <table border="0"> <tr> <td>9816335.5</td> <td>27 July 1998 (27.07.98)</td> <td>GB</td> </tr> <tr> <td>60/125,163</td> <td>19 March 1999 (19.03.99)</td> <td>US</td> </tr> </table> <p>(71) Applicant (for all designated States except US): MICROBIAL TECHNICS LIMITED [GB/GB]; 20 Trumpington Street, Cambridge CB2 1QA (GB).</p> <p>(72) Inventors; and</p> <p>(75) Inventors/Applicants (for US only): LE PAGE, Richard, William, Falla [GB/GB]; University of Cambridge, Dept. of Pathology, Tennis Court Road, Cambridge CB2 1QP (GB). WELLS, Jeremy, Mark [GB/GB]; Institute of Food Re- search, Norwich Laboratory, Norwich Research Park, Col- ney, Norwich NR4 7UA (GB). HANNIFFY, Sean, Bosco [IE/GB]; University of Cambridge, Dept. of Pathology, Ten- nis Court Road, Cambridge CB2 1QP (GB).</p> <p>(74) Agents: CHAPMAN, Paul, William et al.; Kilburn &amp; Strode, 20 Red Lion Street, London WC1R 4PJ (GB).</p>	9816335.5	27 July 1998 (27.07.98)	GB	60/125,163	19 March 1999 (19.03.99)	US	<p>(81) Designated States: CA, CN, JP, US, European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE).</p> <p><b>Published</b> <i>Without international search report and to be republished upon receipt of that report.</i></p>
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<p>(54) Title: NUCLEIC ACIDS AND PROTEINS FROM GROUP B STREPTOCOCCUS</p>							
<p>(57) Abstract</p> <p>Novel protein antigens from Group B <i>Streptococcus</i> are described, together with nucleic acid sequences encoding them. Their use in vaccines and screening methods is also described.</p>							

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**NUCLEIC ACIDS AND PROTEINS FROM GROUP B STREPTOCOCCUS**

The present invention relates to proteins derived from *Streptococcus agalactiae*, nucleic acid molecules encoding such proteins, and the use of the proteins as antigens and/or immunogens and in detection/diagnosis. It also relates to a method for the rapid  
5 screening of bacterial genomes to isolate and characterise bacterial cell envelope associated or secreted proteins.

The *Group B Streptococcus* (GBS) (*Streptococcus agalactiae*) is an encapsulated bacterium which emerged in the 1970s as a major pathogen of humans causing sepsis  
10 and meningitis in neonates as well as adults. The incidence of early onset neonatal infection during the first 5 days of life varies from 0.7 to 3.7 per 1000 live births and causes mortality in about 20% of cases. Between 25-50% of neonates surviving early onset infections frequently suffer neurological sequelae. Late onset neonatal infections occur from 6 days to three months of age at a rate of about 0.5 - 1.0 per 1000 live  
15 births.

There is an established association between the colonisation of the maternal genetic tract by GBS at the time of birth and the risk of neonatal sepsis. In humans it has been established that the rectum may act as a reservoir for GBS. Susceptibility in the  
20 neonate is correlated with the a low concentration or absence of IgG antibodies to the capsular polysaccharides found on GBS causing human disease. In the USA strains isolated from clinical cases usually belong to capsular serotypes Ia, Ib, II, III although serotype V may be of increasing significance. Type VIII GBS is the major cause of neonatal sepsis in Japan.

25 A possible means of prevention involves intra or postpartum administration of antibiotics to the mother but there are concerns that this might lead to the emergence of resistant organisms and in some cases allergic reactions. Vaccination of the adolescent females to induce long lasting maternally derived immunity is one of the  
30 most promising approaches to prevent GBS infections in neonates. The capsular

polysaccharide antigens of these organisms have attracted most attention as with regard to vaccine development. Studies in healthy adult volunteers have shown that serotype Ia, II and III polysaccharides are non-toxic and immunogenic in approximately 65%, 95% and 70% of non-immune adults respectively. One of the problems with using capsule antigens as vaccines is that the response rates vary according to pre-immunisation status and the polysaccharide antigen and not all vaccinees produce adequate levels of IgG antibody as indicated in vaccination studies with GBS polysaccharides in human volunteers.

Some people do not respond despite repeated stimuli. These properties are due to the T-independent nature of polysaccharide antigens. One strategy to enhance the immunogenicity of these vaccines is to enhance the T cell dependent properties of polysaccharides by conjugating them to a protein. The use of polysaccharide conjugates looks promising but there are still unresolved questions concerning the nature of the carrier protein. A conjugate vaccine against GBS would require at least 4 different conjugates to be prepared adding to the cost of a vaccine.

Recent evidence also suggests that bacterial surface proteins may be useful to confer immunity. A protein called Rib which is found on most serotype III strains but rarely on serotypes Ia, Ib or II confers immunity to challenge with Rib expressing GBS in animal models (Stalhammar-Carlemalm *et al.*, *Journal of Experimental Medicine* 177:1593-1603 (1993)). Another surface protein of interest as a component of a vaccine is the alpha antigen of the C proteins which protected vaccinated mice against lethal infection with strains expressing alpha protein. The amount of antigen expressed by GBS strains varies markedly.

Approaches to vaccination against GBS infections which rely on the use of capsular polysaccharides have the disadvantage that response rates are likely to vary considerably according to pre-immunisation status and the particular type of polysaccharide antigen used. Results of trials in human volunteers have indicated that

response rates may only be around 65% for some of the key capsule antigens (Larsson *et al.*, *Infection and Immunity* **64**:3518-3523 (1996)). It is also not clear whether all individuals responding to the vaccine would have adequate levels of polysaccharide specific IgG which can cross the placenta and afford immunity to neonates. By  
5 conjugating a protein carrier to the polysaccharide antigen it may be possible to convert them to T-cell dependent antigens and enhance their immunogenicity.

Preliminary studies with GBS type III polysaccharide-tetanus toxoid conjugate have been encouraging (Baker *et al.*, *Reviews of Infectious Diseases* **7**:458-467 (1985),  
10 Baker *et al.*, *The New England Journal of Medicine* **319**:1180-1185 (1988), Paoletti *et al.*, *Infection and Immunity* **64**:677-679 (1996), Paoletti *et al.*, *Infection and Immunity* **62**:3236-3243 (1994)) but in developed countries the use of tetanus may be disadvantageous since most adults will have been immunised against tetanus within the past five years. Additional boosters with tetanus toxoid may cause adverse  
15 reactions (Boyer., *Current Opinions in Pediatrics* **7**:13-18 (1995)). The polysaccharide conjugate vaccines have the disadvantage of being costly to produce and manufacture in comparison with many other kinds of vaccines. There is also the possible risk of problems caused by the cross reactivity between GBS polysaccharides and sialic acid-containing human glycoproteins.

20

An alternative to polysaccharides as antigens is the use of protein antigens derived from GBS. Recent evidence suggest that the GBS surface associated proteins Rib and alpha C protein may be used to confer immunity to GBS infections in experimental model systems (Stalhammar-Carlemalm *et al.*, (1993) [*supra*], Larsson *et al.*, (1996)  
25 [*supra*]). However these two proteins are not conserved in all serotypes of GBS which cause disease in humans. Assuming that these antigens would be immunogenic and elicit protective level responses in humans they would not confer protection against all infections as 10% of infectious *Group B streptococci* do not express Rib or C protein  
alpha.

30

This invention seeks to overcome the problem of vaccination against GBS by using a novel screening method specifically designed to identify those *Group B Streptococcus* genes encoding bacterial cell surface associated or secreted proteins (antigens). The proteins expressed by these genes may be immunogenic, and therefore may be useful  
5 in the prevention and treatment of *Group B Streptococcus* infection. For the purposes of this application, the term immunogenic means that these proteins will elicit a protective immune response within a subject. Using this novel screening method a number of genes encoding novel *Group B Streptococcus* proteins have been identified.

10 Thus in a first aspect, the present invention provides a *Group B Streptococcus* protein, having a sequence selected from those shown in figure 1, or fragments or derivatives thereof.

It will be apparent to the skilled person that proteins and polypeptides included within  
15 this group may be cell surface receptors, adhesion molecules, transport proteins, membrane structural proteins, and/or signalling molecules.

Alterations in the amino acid sequence of a protein can occur which do not affect the function of a protein. These include amino acid deletions, insertions and substitutions  
20 and can result from alternative splicing and/or the presence of multiple translation start sites and stop sites. Polymorphisms may arise as a result of the infidelity of the translation process. Thus changes in amino acid sequence may be tolerated which do not affect the proteins function.

25 Thus, the present invention includes derivatives or variants of the proteins, polypeptides, and peptides of the present invention which show at least 50% identity to the proteins, polypeptides and peptides described herein. Preferably the degree of sequence identity is at least 60% and preferably it is above 75%. More preferably still is it above 80%, 90% or even 95%.

30

The term identity can be used to describe the similarity between two polypeptide sequences. A software package well known in the art for carrying out this procedure is the CLUSTAL program. It compares the amino acid sequences of two polypeptides and finds the optimal alignment by inserting spaces in either sequence as appropriate.

5 The amino acid identity or similarity (identity plus conservation of amino acid type) for an optimal alignment can also be calculated using a software package such as BLASTx. This program aligns the largest stretch of similar sequence and assigns a value to the fit. For any one pattern comparison several regions of similarity may be found, each having a different score. One skilled in the art will appreciate that two  
10 polypeptides of different lengths may be compared over the entire length of the longer fragment. Alternatively small regions may be compared. Normally sequences of the same length are compared for a useful comparison to be made.

15 Manipulation of the DNA encoding the protein is a particularly powerful technique for both modifying proteins and for generating large quantities of protein for purification purposes. This may involve the use of PCR techniques to amplify a desired nucleic acid sequence. Thus the sequence data provided herein can be used to design primers for use in PCR so that a desired sequence can be targeted and then amplified to a high degree.

20 Typically primers will be at least five nucleotides long and will generally be at least ten nucleotides long (e.g. fifteen to twenty-five nucleotides long). In some cases primers of at least thirty or at least thirty-five nucleotides in length may be used.

25 As a further alternative chemical synthesis may be used. This may be automated. Relatively short sequences may be chemically synthesised and ligated together to provide a longer sequence.

Thus in a further aspect, the present invention provides , a nucleic acid molecule comprising or consisting of a sequence which is:

- (i) any of the DNA sequences set out in figure 1 herein or their RNA equivalents;
- (ii) a sequence which is complementary to any of the sequences of (i);
- (iii) a sequence which codes for the same protein or polypeptide, as those sequences of (i) or (ii);
- (iv) a sequence which shows substantial identity with any of those of (i), (ii) and (iii); or
- (v) a sequence which codes for a derivative or fragment of a nucleic acid molecule shown in figure 1.

The term identity can also be used to describe the similarity between two individual DNA sequences. The 'bestfit' program (Smith and Waterman, *Advances in applied Mathematics*, 482-489 (1981)) is one example of a type of computer software used to find the best segment of similarity between two nucleic acid sequences, whilst the GAP program enables sequences to be aligned along their whole length and finds the optimal alignment by inserting spaces in either sequence as appropriate.

The term 'RNA equivalent' when used above indicates that a given RNA molecule has a sequence which is complementary to that of a given DNA molecule, allowing for the fact that in RNA 'U' replaces 'T' in the genetic code. The nucleic acid molecule may be in isolated or recombinant form.

The nucleic acid molecule may be in an isolated or recombinant form. DNA constructs can readily be generated using methods well known in the art. These techniques are disclosed, for example in J. Sambrook *et al*, *Molecular Cloning 2<sup>nd</sup> Edition*, Cold Spring Harbour Laboratory Press (1989). Modifications of DNA constructs and the proteins expressed such as the addition of promoters, enhancers, signal sequences, leader sequences, translation start and stop signals and DNA stability controlling regions, or the addition of fusion partners may then be facilitated.



Normally the DNA construct will be inserted into a vector which may be of phage or plasmid origin. Expression of the protein is achieved by the transformation or transfection of the vector into a host cell which may be of eukaryotic or prokaryotic origin. Such vectors and suitable host cells form yet further aspects of the present invention.

The *Group B Streptococcus* proteins (antigens) described herein can additionally be used to raise antibodies, or to generate affibodies. These can be used to detect *Group B Streptococcus*.

Thus in a further aspect the present invention provides, an antibody, affibody, or a derivative thereof which binds to any one or more of the proteins, polypeptides, peptides, fragments or derivatives thereof, as described herein.

Antibodies within the scope of the present invention may be monoclonal or polyclonal. Polyclonal antibodies can be raised by stimulating their production in a suitable animal host (e.g. a mouse, rat, guinea pig, rabbit, sheep, goat or monkey) when a protein as described herein, or a homologue, derivative or fragment thereof, is injected into the animal. If desired, an adjuvant may be administered together with the protein. Well-known adjuvants include Freund's adjuvant (complete and incomplete) and aluminium hydroxide. The antibodies can then be purified by virtue of their binding to a protein as described herein.

Monoclonal antibodies can be produced from hybridomas. These can be formed by fusing myeloma cells and spleen cells which produce the desired antibody in order to form an immortal cell line. Thus the well-known Kohler & Milstein technique (*Nature* **256** (1975)) or subsequent variations upon this technique can be used.

Techniques for producing monoclonal and polyclonal antibodies that bind to a particular polypeptide/protein are now well developed in the art. They are discussed in standard

immunology textbooks, for example in Roitt *et al*, *Immunology* second edition (1989), Churchill Livingstone, London.

5 In addition to whole antibodies, the present invention includes derivatives thereof which are capable of binding to proteins etc as described herein. Thus the present invention includes antibody fragments and synthetic constructs. Examples of antibody fragments and synthetic constructs are given by Dougall *et al* in *Tibtech* **12** 372-379 (September 1994).

10 Antibody fragments include, for example, Fab, F(ab')<sub>2</sub> and Fv fragments. Fab fragments (These are discussed in Roitt *et al* [*supra*]). Fv fragments can be modified to produce a synthetic construct known as a single chain Fv (scFv) molecule. This includes a peptide linker covalently joining V<sub>h</sub> and V<sub>l</sub> regions, which contributes to the stability of the molecule. Other synthetic constructs that can be used include CDR peptides. These are  
15 synthetic peptides comprising antigen-binding determinants. Peptide mimetics may also be used. These molecules are usually conformationally restricted organic rings that mimic the structure of a CDR loop and that include antigen-interactive side chains.

Synthetic constructs include chimaeric molecules. Thus, for example, humanised (or  
20 primatised) antibodies or derivatives thereof are within the scope of the present invention. An example of a humanised antibody is an antibody having human framework regions, but rodent hypervariable regions. Ways of producing chimaeric antibodies are discussed for example by Morrison *et al* in *PNAS*, **81**, 6851-6855 (1984) and by Takeda *et al* in *Nature*. **314**, 452-454 (1985).

25 Synthetic constructs also include molecules comprising an additional moiety that provides the molecule with some desirable property in addition to antigen binding. For example the moiety may be a label (e.g. a fluorescent or radioactive label). Alternatively, it may be a pharmaceutically active agent.

30

Affibodies are proteins which are found to bind to target proteins with a low dissociation constant. They are selected from phage display libraries expressing a segment of the target protein of interest (Nord K, Gunneriusson E, Ringdahl J, Stahl S, Uhlen M, Nygren PA, Department of Biochemistry and Biotechnology, Royal Institute of Technology (KTH), Stockholm, Sweden).

In a further aspect the invention provides an immunogenic composition comprising one or more proteins, polypeptides, peptides, fragments or derivatives thereof, or nucleotide sequences described herein. A composition of this sort may be useful in the treatment or prevention of *Group B Streptococcus* infection in subject. In a preferred aspect of the invention the immunogenic composition is a vaccine.

In other aspects the invention provides:

- i) Use of an immunogenic composition as described herein in the preparation of a medicament for the treatment or prophylaxis of *Group B Streptococcus* infection. Preferably the medicament is a vaccine.
- ii) A method of detection of *Group B Streptococcus* which comprises the step of bringing into contact a sample to be tested with at least one antibody, affibody, or a derivative thereof, as described herein.
- iii) A method of detection of *Group B Streptococcus* which comprises the step of bringing into contact a sample to be tested with at least one protein, polypeptide, peptide, fragments or derivatives as described herein.
- iv) A method of detection of *Group B Streptococcus* which comprises the step of bringing into contact a sample to be tested with at least one nucleic acid molecule as described herein.

- v) A kit for the detection of *Group B Streptococcus* comprising at least one antibody, affibody, or derivatives thereof, described herein.
- 5 vi) A kit for the detection of *Group B Streptococcus* comprising at least one *Group B Streptococcus* protein, polypeptide, peptide, fragment or derivative thereof, as described herein.
- vii) A kit for the detection of *Group B Streptococcus* comprising at least one nucleic acid of the invention.

10

As described previously, the novel proteins described herein are identified and isolated using a novel screening method which specifically identifies those *Group B Streptococcus* genes encoding bacterial cell envelope associated or secreted proteins.

15

The information necessary for the secretion/export of proteins has been extensively studied in bacteria. In the majority of cases, export requires a signal peptide positioned at the N-terminus of the precursor protein to target the precursor to translocation sites on the membrane. During or after translocation, the signal peptide is removed by a signal peptidase. The ultimate destination/localisation of the protein, (whether it be

20 secreted extracellularly or anchored to the bacterium's surface, etc) is determined by sequences other than the leader peptide sequence.

25

Recently, Poquet *et al.* (*J. Bacteriol.* **180**:1904-1912 (1998)) have described a screening vector incorporating the *nuc* gene lacking its own signal leader as a reporter to identify exported proteins in Gram positive bacteria, and have applied it to *L. lactis*. Staphylococcal nuclease is a naturally secreted heat-stable, monomeric enzyme which has been efficiently expressed and secreted in a range of Gram positive bacteria (Shortle., *Gene* **22**:181-189 (1983), Kovacevic *et al.*, *J. Bacteriol.* **162**:521-528 (1985), Miller *et al.*, *J. Bacteriol.* **169**:3508-3514 (1987), Liebl *et al.*, *J. Bacteriol.*

174:1854-1861(1992), Le Loir *et al.*, *J. Bacteriol.* **176**:5135-5139 (1994), Poquet *et al.*, 1998 [*supra*]). The screening vector (pFUN) contains the pAM $\beta$ 1 replicon which functions in a broad host range of Gram-positive bacteria in addition to the ColE1 replicon that promotes replication in *Escherichia coli* and certain other Gram  
5 negative bacteria. Unique cloning sites present in the vector can be used to generate transcriptional and translational fusions between cloned genomic DNA fragments and the open reading frame of the truncated *nuc* gene devoid of its own signal secretion leader. The *nuc* gene makes an ideal reporter gene because the secretion of nuclease can readily be detected using a simple and sensitive plate test: Recombinant colonies  
10 secreting the nuclease develop a pink halo whereas control colonies remain white (Shortle, 1983 [*supra*], Le Loir *et al.*, 1994 [*supra*]).

A direct screen to identify and isolate DNA encoding bacterial cell envelope associated or secreted proteins (antigens) in pathogenic bacteria has been developed by  
15 the present inventors which utilises a vector-system (pTREP1 expression vector) in *Lactococcus lactis* that specifically detects DNA sequences which are adjacent to, and associated with DNA encoding surface proteins from *Group B Streptococcus*. The screening vector also incorporates the *nuc* gene encoding the *Staphylococcal* nuclease as a reporter gene.

20 Only the part of the *nuc* gene encoding the mature nuclease protein (minus its signal peptide sequence) is cloned into the pTREP1 expression vector in *L. lactis*. In this form, the *nuc*-encoded nuclease cannot be secreted even when expressed intracellularly. The reporter vector is then randomly combined with appropriately  
25 digested genomic DNA from *Group B Streptococcus*, cloned into *L. lactis* and used as a screening system for sequences permitting the export of nuclease. In this way gene/partial gene sequences encoding exported proteins from *Group B Streptococcus* are isolated. Once a partial gene sequence is obtained, full length sequences encoding exported proteins can readily be obtained using techniques well known in the art.

In possessing a promoter, the pTREP1-*nuc* vectors differ from the pFUN vector described by Poquet *et al.* (1998) [*supra*], which was used to identify *L. lactis* exported proteins by screening directly for *Nuc* activity directly in *L. lactis*. As the  
5 pFUN vector does not contain a promoter upstream of the *nuc* open reading frame the cloned genomic DNA fragment must also provide the signals for transcription in addition to those elements required for translation initiation and secretion of *Nuc*. This limitation may prevent the isolation of genes that are distant from a promoter for example genes which are within polycistronic operons. Additionally there can be no  
10 guarantee that promoters derived from other species of bacteria will be recognised and functional in *L. lactis*. Certain promoters may be under stringent regulation in the natural host but not in *L. lactis*. In contrast, the presence of the P1 promoter in the pTREP1-*nuc* series of vectors ensures that promoterless DNA fragments (or DNA fragments containing promoter sequences not active in *L. lactis*) may still be  
15 transcribed. Thus yet another advantage of this invention is that genes missed in other screening methods may be identified.

Hence in a further aspect the present invention provides a method of screening for DNA encoding bacterial cell wall associated or surface antigens in gram positive  
20 bacteria comprising the steps of:

- combining a reporter vector including the nucleotide sequence encoding the mature form of the staphylococcus nuclease gene and an upstream promoter region with DNA from a gram positive bacteria.
- transforming the resultant vector into *Lactococcus lactis* cells.
- 25 - assaying for the secretion of *staphylococcus* nuclease protein in the transformed cells.

Preferably, the reporter vector is one of the pTREP1-*nuc* vectors shown in figure 4.

In another aspect, the present invention provides a vector as shown in figure 4 for use in screening for DNA encoding exported or surface antigens in gram positive bacteria. Examples of gram positive bacteria which may be screened include *Group B Streptococcus*, *Streptococcus pneumoniae*, *Staphylococcus aureus* or pathogenic  
5 *Group A Streptococci*.

Given that the inventors have identified a group of important proteins, such proteins are potential targets for anti-microbial therapy. It is necessary, however, to determine whether each individual protein is essential for the organism's viability.  
10 Thus, the present invention also provides a method of determining whether a protein or polypeptide as described herein represents a potential anti-microbial target which comprises inactivating said protein and determining whether *Group B Streptococcus* is still viable.

15 A suitable method for inactivating the protein is to effect selected gene knockouts, ie prevent expression of the protein and determine whether this results in a lethal change. Suitable methods for carrying out such gene knockouts are described in Li *et al*, *P.N.A.S.*, **94**:13251-13256 (1997) and Kolkman *et al*

20 In a final aspect the present invention provides the use of an agent capable of antagonising, inhibiting or otherwise interfering with the function or expression of a protein or polypeptide of the invention in the manufacture of a medicament for use in the treatment or prophylaxis of *Group B Streptococcus* infection.

25 The invention will now be described by means of the following example which should not in any way be construed as limiting. The examples refer to the figures in which

Fig 1: (A) Shows a number of full length nucleotide sequences encoding antigenic *Group B Streptococcus* proteins. (B) Shows the corresponding amino acid sequences.

5 Fig 2: Shows a number of oligonucleotide primers used in the screening process

**nucS1** primer designed to amplify a mature form of the nuc A gene

**nucS2-** primer designed to amplify a mature form of the nuc A gene.

**nucS3** primer designed to amplify a mature form of the nuc A gene

10 **nucR** primer designed to amplify a mature form of the nuc A gene

**nucseq** primer designed to sequence DNA cloned into the pTREP-Nuc vector

**pTREPF** nucleic acid sequence containing recognition site for ECORV. Used for cloning fragments into pTREX7.

15 **pTREPR** nucleic acid sequence containing recognition site for BAMH1. Used for cloning fragments into pTREX7.

**PUCF** forward sequencing primer, enables direct sequencing of cloned DNA fragments.

**VR** example of gene specific primer used to obtain further antigen DNA sequence by the method of DNA walking.

20 **V1** example of gene specific primer used to obtain further antigen DNA sequence by the method of DNA walking.

**V2** example of gene specific primer used to obtain further antigen DNA sequence by the method of DNA walking.

25

Fig 3: (i) Schematic presentation of the nucleotide sequence of the unique gene cloning site immediately upstream of the mature *nuc* gene in pTREP1-*nuc1*, pTREP1-*nuc2* and pTREP1-*nuc3*. Each of the pTREP-*nuc* vectors contain an



EcoRV (a SmaI site in pTREP1-*nuc2*) cleavage site which allows cloning of genomic DNA fragments in 3 different frames with respect to the mature *nuc* gene.

(ii) A physical and genetic summary map of the pTREP1-*nuc* vectors. The expression cassette incorporating *nuc*, the macrolides, lincosamides and streptogramin B (MLS) resistance determinant, and the replicon (rep) *Ori-pAMβ1* are depicted (not drawn to scale).

(iii) Schematic presentation of the expression cassette showing the various sequence elements involved in gene expression and location of unique restriction endonuclease sites (not drawn to scale).

Fig 4: Shows the results of various DNA vaccine trials;

Fig 5: Shows the results of a second group of DNA vaccine trials;

Figs 6-11: Show various Southern Blot analyses of different Group B streptococcus strains.

### Example 1

Thus far more than 100 gene/partial gene sequences putatively encoding exported proteins in *S. agalactiae* have been identified using the nuclease screening system of the invention. These have been further analysed to remove artifacts. The nucleotide sequences of genes identified using the screening system has been characterised using a number of parameters described below. All of these sequences are novel in that they have not been described previously.

1. All putative surface proteins are analysed for leader/signal peptide sequences. Bacterial signal peptide sequences share a common design. They are characterised by a short positively charged N-terminus (N region) immediately preceding a stretch of hydrophobic residues (central portion-h region) followed by a

more polar C-terminal portion which contains the cleavage site (c-region). Computer software is used to perform hydropathy profiling of putative proteins (Marcks, *Nuc. Acid. Res.*, **16**:1829-1836 (1988)) which is used to identify the distinctive hydrophobic portion (h-region) typical of leader peptide sequences. In addition, the presence/absence of a potential ribosomal binding site (Shine-Dalgarno sequence required for translation) is also noted.

2. All putative surface protein sequences are used to search the OWL sequence database which includes a translation of the GENBANK and SWISSPROT database.. This allows identification of similar sequences which may have been previously characterised not only at the sequence level but at a functional level. It may also provide information indicating that these proteins are indeed surface related and not artifacts.

3. Putative *S. agalactiae* surface proteins are also be assessed for their novelty. Some of the identified proteins may or may not possess a typical leader peptide sequence and may not show homology with any DNA/protein sequences in the database. Indeed these proteins may indicate the primary advantage of our screening method, i.e. isolating atypical surface-related proteins, which would have been missed in all previously described screening protocols.

The construction of three reporter vectors and their use in *L. lactis* to identify and isolate genomic DNA fragments from pathogenic bacteria encoding secreted or surface associated proteins is now described.

## **Construction of the pTREP1-*nuc* series of reporter vectors**

### **(a) Construction of expression plasmid pTREP1**

The pTREP1 plasmid is a high-copy number (40-80 per cell) theta-replicating gram positive plasmid, which is a derivative of the pTREX plasmid which is itself a derivative of the the previously published pIL253 plasmid. pIL253 incorporates the

broad Gram-positive host range replicon of pAM $\beta$ 1 (Simon and Chopin, 1988) and is non-mobilisable by the *L lactis* sex-factor. pIL253 also lacks the *tra* function which is necessary for transfer or efficient mobilisation by conjugative parent plasmids exemplified by pIL501. The Enterococcal pAM $\beta$ 1 replicon has previously been transferred to various species including *Streptococcus*, *Lactobacillus* and *Bacillus* species as well as *Clostridium acetobutylicum*, (LeBlanc *et al.*, *Proceedings of the National Academy of Science USA* **75**:3484-3487 (1978)) indicating the potential broad host range utility. The pTREP1 plasmid represents a constitutive transcription vector.

The pTREX vector was constructed as follows. An artificial DNA fragment containing a putative RNA stabilising sequence, a translation initiation region (TIR), a multiple cloning site for insertion of the target genes and a transcription terminator was created by annealing 2 complementary oligonucleotides and extending with Tfl DNA polymerase. The sense and anti-sense oligonucleotides contained the recognition sites for NheI and BamHI at their 5' ends respectively to facilitate cloning. This fragment was cloned between the XbaI and BamHI sites in pUC19NT7, a derivative of pUC19 which contains the T7 expression cassette from pLET1 (Wells *et al.*, *J. Appl. Bacteriol.* **74**:629-636 (1993)) cloned between the EcoRI and HindIII sites. The resulting construct was designated pUCLEX. The complete expression cassette of pUCLEX was then removed by cutting with HindIII and blunting followed by cutting with EcoRI before cloning into EcoRI and SacI (blunted) sites of pIL253 to generate the vector pTREX (Wells and Schofield, *In Current advances in metabolism, genetics and applications-NATO ASI Series. H* **98**:37-62. (1996)). The putative RNA stabilising sequence and TIR are derived from the *Escherichia coli* T7 bacteriophage sequence and modified at one nucleotide position to enhance the complementarity of the Shine Dalgarno (SD) motif to the ribosomal 16s RNA of *Lactococcus lactis* (Schofield *et al.* pers. coms. University of Cambridge Dept. Pathology.).

A *Lactococcus lactis* MG1363 chromosomal DNA fragment exhibiting promoter activity which was subsequently designated P7 was cloned between the EcoRI and BglII sites present in the expression cassette, creating pTREX7. This active promoter region had been previously isolated using the promoter probe vector pSB292  
5 (Waterfield *et al.*, *Gene* **165**:9-15 (1995)). The promoter fragment was amplified by PCR using the Vent DNA polymerase according to the manufacturer.

The pTREP1 vector was then constructed as follows. An artificial DNA fragment which included a transcription terminator, the forward pUC sequencing primer, a  
10 promoter multiple cloning site region and a universal translation stop sequence was created by annealing two overlapping partially complementary synthetic oligonucleotides together and extending with sequenase according to manufacturers instructions. The sense and anti-sense (pTREP<sub>F</sub> and pTREP<sub>R</sub>) oligonucleotides contained the recognition sites for EcoRV and BamHI at their 5' ends respectively to  
15 facilitate cloning into pTREX7. The transcription terminator was that of the *Bacillus penicillinase* gene, which has been shown to be effective in *Lactococcus* (Jos *et al.*, *Applied and Environmental Microbiology* **50**:540-542 (1985)). This was considered necessary as expression of target genes in the pTREX vectors was observed to be leaky and is thought to be the result of cryptic promoter activity in the origin region  
20 (Schofield *et al.* pers. coms. University of Cambridge Dept. Pathology.). The forward pUC primer sequencing was included to enable direct sequencing of cloned DNA fragments. The translation stop sequence which encodes a stop codon in 3 different frames was included to prevent translational fusions between vector genes and cloned DNA fragments. The pTREX7 vector was first digested with EcoRI and blunted using  
25 the 5' - 3' polymerase activity of T4 DNA polymerase (NEB) according to manufacturer's instructions. The EcoRI digested and blunt ended pTREX7 vector was then digested with Bgl II thus removing the P7 promoter. The artificial DNA fragment derived from the annealed synthetic oligonucleotides was then digested with EcoRV and Bam HI and cloned into the EcoRI(blunted)-Bgl II digested pTREX7 vector to

generate pTREP. A *Lactococcus lactis* MG1363 chromosomal promoter designated P1 was then cloned between the EcoRI and BglII sites present in the pTREP expression cassette forming pTREP1. This promoter was also isolated using the promoter probe vector pSB292 and characterised by Waterfield *et al.*, (1995) [*supra*]. The P1 promoter fragment was originally amplified by PCR using vent DNA polymerase according to manufacturers instructions and cloned into the pTREP as an EcoRI-BglII DNA fragment. The EcoRI-BglII P1 promoter containing fragment was removed from pTREP1 by restriction enzyme digestion and used for cloning into pTREP (Schofield *et al.* pers. coms. University of Cambridge, Dept. Pathology.).

**(b) PCR amplification of the *S. aureus nuc* gene.**

The nucleotide sequence of the *S. aureus nuc* gene (EMBL database accession number V01281) was used to design synthetic oligonucleotide primers for PCR amplification. The primers were designed to amplify the mature form of the *nuc* gene designated *nucA* which is generated by proteolytic cleavage of the N-terminal 19 to 21 amino acids of the secreted propeptide designated Snase B (Shortle, 1983 [*supra*]). Three sense primers (*nucS1*, *nucS2* and *nucS3*, shown in figure 3) were designed, each one having a blunt-ended restriction endonuclease cleavage site for EcoRV or SmaI in a different reading frame with respect to the *nuc* gene. Additionally BglII and BamHI were incorporated at the 5' ends of the sense and anti-sense primers respectively to facilitate cloning into BamHI and BglII cut pTREP1. The sequences of all the primers are given in figure 3. Three *nuc* gene DNA fragments encoding the mature form of the nuclease gene (*NucA*) were amplified by PCR using each of the sense primers combined with the anti-sense primer. The *nuc* gene fragments were amplified by PCR using *S. aureus* genomic DNA template, Vent DNA Polymerase (NEB) and the conditions recommended by the manufacturer. An initial denaturation step at 93°C for 2 min was followed by 30 cycles of denaturation at 93°C for 45 sec, annealing at 50°C for 45 seconds, and extension 73°C for 1 minute and then a final 5 min extension step

at 73°C. The PCR amplified products were purified using a Wizard clean up column (Promega) to remove unincorporated nucleotides and primers.

**(c) Construction of the pTREP1-*nuc* vectors**

5

The purified *nuc* gene fragments described in section b were digested with Bgl II and BamHI using standard conditions and ligated to BamHI and BglIII cut and dephosphorylated pTREP1 to generate the pTREP1-*nuc*1, pTREP1-*nuc*2 and pTREP1-*nuc*3 series of reporter vectors. These vectors are described in figure 4.

10

General molecular biology techniques were carried out using the reagents and buffers supplied by the manufacturer or using standard techniques (Sambrook and Maniatis, Molecular cloning: A laboratory manual. Cold Spring Harbor Laboratory Press: Cold Spring Harbour (1989)). In each of the pTREP1-*nuc* vectors the expression cassette

15

comprises a transcription terminator, lactococcal promoter P1, unique cloning sites (BglIII, EcoRV or SmaI) followed by the mature form of the *nuc* gene and a second transcription terminator. Note that the sequences required for translation and secretion of the *nuc* gene were deliberately excluded in this construction. Such elements can only be provided by appropriately digested foreign DNA fragments (representing the target bacterium) which can be cloned into the unique restriction sites present immediately upstream of the *nuc* gene.

20

**(d) Screening for secreted proteins in *Group B Streptococcus*.**

Genomic DNA isolated from and *Group B Streptococcus* (*S. agalactiae*) was digested with the restriction enzyme Tru9I. This enzyme which recognises the sequence 5'-TTAA -3' was used because it cuts A/T rich genomes efficiently and can generate random genomic DNA fragments within the preferred size range (usually averaging 0.5 - 1.0 kb). This size range was preferred because there is an increased probability that the P1 promoter can be utilised to transcribe a novel gene sequence. However, the P1 promoter may not be necessary in all cases as it is possible that many Streptococcal promoters are recognised in *L. lactis*. DNA fragments of different size ranges were

25

30

purified from partial Tru9I digests of and *S. agalactiae* genomic DNA. As the Tru 9I restriction enzyme generates staggered ends the DNA fragments had to be made blunt ended before ligation to the EcoRV or SmaI cut pTREP1-*nuc* vectors. This was achieved by the partial fill-in enzyme reaction using the 5'-3' polymerase activity of Klenow enzyme. Briefly Tru9I digested DNA was dissolved in a solution (usually between 10-20  $\mu$ l in total) supplemented with T4 DNA ligase buffer (New England Biolabs; NEB) (1X) and 33  $\mu$ M of each of the required dNTPs, in this case dATP and dTTP. Klenow enzyme was added (1 unit Klenow enzyme (NEB) per  $\mu$ g of DNA) and the reaction incubated at 25°C for 15 minutes. The reaction was stopped by incubating the mix at 75°C for 20 minutes. EcoRV or SmaI digested pTREP-*nuc* plasmid DNA was then added (usually between 200-400 ng). The mix was then supplemented with 400 units of T4 DNA ligase (NEB) and T4 DNA ligase buffer (1X) and incubated overnight at 16°C. The ligation mix was precipitated directly in 100% Ethanol and 1/10 volume of 3M sodium acetate (pH 5.2) and used to transform *L. lactis* MG1363 (Gasson, *J. Bacteriol.* **154**:1-9 (1983)). Alternatively, the gene cloning site of the pTREP-*nuc* vectors also contains a BglIII site which can be used to clone for example Sau3AI digested genomic DNA fragments.

*L. lactis* transformant colonies were grown on brain heart infusion agar and nuclease secreting (*Nuc*<sup>+</sup>) clones were detected by a toluidine blue-DNA-agar overlay (0.05 M Tris pH 9.0, 10 g of agar per litre, 10 g of NaCl per liter, 0.1 mM CaCl<sub>2</sub>, 0.03% wt/vol. salmon sperm DNA and 90 mg of Toluidine blue O dye) essentially as described by Shortle, 1983 [*supra*], and Le Loir *et al.*, 1994 [*supra*]). The plates were then incubated at 37°C for up to 2 hours. Nuclease secreting clones develop an easily identifiable pink halo. Plasmid DNA was isolated from *Nuc*<sup>+</sup> recombinant *L. lactis* clones and DNA inserts were sequenced on one strand using the *NucSeq* sequencing primer described in figure 3, which sequences directly through the DNA insert.

Whilst the example described above related specifically to *Group B Streptococcus*, it will be apparent to one skilled in the art that the same screening technique may be used to detect exported and secreted proteins in other gram positive bacteria, for example *Streptococcus pneumoniae*.

5 **Example 2; Screening Group B Streptococcal derived genes in DNA vaccination experiments.**

**pcDNA3.1+ as a DNA vaccine vector**

10 The commercially available pcDNA3.1+ plasmid (Invitrogen), referred to as pcDNA3.1 henceforth, was used as a vector in all DNA immunisation experiments involving gene targets derived using the LEEP system. pcDNA 3.1 is designed for high-level stable and transient expression in mammalian cells and has been used widely and successfully as a host vector to test candidate genes from a variety of pathogens in DNA vaccination experiments (Zhang *et al.*, 1997; Kurar and Splitter, 15 1997; Anderson *et al.*, 1996).

The vector possesses a multiple cloning site which facilitates the cloning of multiple gene targets downstream of the human cytomegalovirus (CMV) immediate-early promoter/enhancer which permits efficient, high-level expression of the target gene in 20 a wide variety of mammalian cells and cell types including both muscle and immune cells. This is important for optimal immune response as it remains unknown as to which cells types are most important in generating a protective response *in vivo*. The plasmid also contains the ColE1 origin of replication which allows convenient high-copy number replication and growth in *E. coli* and the ampicillin resistance gene (B-lactamase) for selection in *E. coli*. In addition pcDNA 3.1 possesses a T7 25 promoter/priming site upstream of the MCS which allows for *in vitro* transcription of a cloned gene in the sense orientation.

30 **Preparation of DNA vaccines**

Oligonucleotide primers were designed for each individual gene of interest derived using the LEEP system. Each gene was examined thoroughly, and where possible, primers were designed such that they targeted that portion of the gene thought to



5 encode only the mature portion of the protein (**APPENDIX I**). It was hoped that  
expressing those sequences that encode only the mature portion of a target gene  
protein, would facilitate its correct folding when expressed in mammalian cells. For  
example, in the majority of cases primers were designed such that putative N-terminal  
10 signal peptide sequences would not be included in the final amplification product to be  
cloned into the pcDNA3.1 expression vector. The signal peptide directs the  
polypeptide precursor to the cell membrane via the protein export pathway where it is  
normally cleaved off by signal peptidase I (or signal peptidase II if a lipoprotein).  
Hence the signal peptide does not make up any part of the mature protein whether it be  
15 displayed on the bacterium's surface or secreted. Where a N-terminal leader peptide  
sequence was not immediately obvious, primers were designed to target the whole of  
the gene sequence for cloning and ultimately, expression in pcDNA3.1.

15 All forward and reverse oligonucleotide primers incorporated appropriate restriction  
enzyme sites to facilitate cloning into the pcDNA3.1 MCS region. All forward primers  
were also designed to include the conserved Kozak nucleotide sequence 5'-gccacc-3'  
immediately upstream of an 'atg' translation initiation codon in frame with the target  
gene insert. The Kozak sequence facilitates the recognition of initiator sequences by  
eukaryotic ribosomes. Typically, a forward primer incorporating a BamH1 restriction  
20 enzyme site the primer would begin with the sequence 5'-cgggatccgccaccatg-3',  
followed by a sequence homologous to the 5' end of that part of a gene being  
amplified. All reverse primers incorporated a Not I restriction enzyme site sequence 5'  
-ttcgggccgc-3'. All gene-specific forward and reverse primers were designed with  
compatible melting temperatures to facilitate their amplification.

25 All gene targets were amplified by PCR from *S. agalactiae* genomic DNA template  
using Vent DNA polymerase (NEB) or r*Tth* DNA polymerase (*PE Applied*  
*Biosystems*) using conditions recommended by the manufacturer. A typical  
amplification reaction involved an initial denaturation step at 95°C for 2 minutes  
30 followed by 35 cycles of denaturation at 95°C for 30 seconds, annealing at the  
appropriate melting temperature for 30 seconds, and extension at 72°C for 1 minute (1  
minute per kilobase of DNA being amplified). This was followed by a final extension  
period at 72°C for 10 minutes. All PCR amplified products were extracted once with  
phenol chloroform (2:1:1) and once with chloroform (1:1) and ethanol precipitated.

Specific DNA fragments were isolated from agarose gels using the QIAquick Gel Extraction Kit (Qiagen). The purified amplification gene DNA fragments were digested with the appropriate restriction enzymes and cloned into the pcDNA3.1 plasmid vector using *E. coli* as a host. Successful cloning and maintenance of genes was confirmed by restriction mapping and by DNA sequencing. Recombinant plasmid DNA was isolated on a large scale (>1.5 mg) using Plasmid Mega Kits (Qiagen).

It was decided to include the *S. agalactiae rib* gene as a positive control in at least one trial of DNA immunisation experiments. Rabbit antiserum against the Rib protein (Stalhammar-Carleman *et al.*, 1993) and highly purified preparations of the Rib protein itself (Larsson *et al.*, 1999; Larsson *et al.*, 1996) have been shown to confer protection against lethal infection with strains expressing the antigen. All serotype III strains have been shown to express the Rib antigen and Southern blot analysis performed in the laboratory has confirmed that *S. agalactiae* serotype III (strain 97/0099) does contain the *rib* gene, hence the *rib* gene as part of a DNA vaccine would represent a potential positive control for all DNA immunisation experiments. Oligonucleotide primers were designed (**Appendix I**) that targeted only the mature portion of the *rib* gene and which included appropriate restriction enzyme sites for cloning into pcDNA3.1. *rib* was amplified using rTth DNA polymerase (*PE Applied Biosystems*) using conditions recommended by the manufacturer. Conditions for cloning were similar to that described previously.

### **Preparation of a *S. agalactiae* standard inoculum**

#### **Strain validation**

*S. agalactiae* serotype III (strain 97/0099) is a recent clinical isolate derived from the cerebral spinal fluid of a new born baby suffering from meningitis. This haemolytic strain of Group B Streptococcus was epidemiologically tested and validated at the Respiratory and Systemic Infection Laboratory, PHLS Central Public health laboratory, 61 Collindale Avenue, London NW9 5HT. The strain was subcultured only twice prior to its arrival in the laboratory. Upon its arrival on a agar slope, a sweep of 4-5 colonies was immediately used to inoculate a Todd Hewitt/5% horse blood broth which was incubated overnight statically at 37 C. 0.5 ml aliquots of this overnight culture were then used to make 20% glycerol stocks of the bacterium for long term

storage at -70 °C. Glycerol stocks were streaked on Todd Hewitt/5% horse blood agar plates to confirm viability.

#### 5 ***In vivo* passaging of Group B Streptococcus**

A frozen culture (described under strain validation) of *S. agalactiae* serotype III (strain 97/0099) was streaked to single colonies on Todd-Hewitt/5% blood agar plates which were incubated overnight at 37°C. A sweep of 4-5 colonies was used to inoculate a Todd Hewitt/5% horse blood broth which was again incubated overnight. A 0.5 ml  
10 aliquot from this overnight culture was used to inoculate a 50 ml Todd Hewitt broth (1:100 dilution) which was incubated at 37 °C. 10-fold serial dilutions of the overnight culture were made (since virulence of this strain was unknown) and each were passaged intra-peritoneally (IP) in CBA/ca mice in duplicate. Viable counts were performed on the various inocula used in the passage. Groups of mice were challenged  
15 with various concentrations of the pathogen ranging from 10<sup>8</sup> to 10<sup>4</sup> colony forming units (cfu). Mice that developed symptoms were terminally anaesthetized and cardiac punctures were performed (Only mice that had been challenged with the highest doses, i.e. 1 X 10<sup>8</sup> cfu, developed symptoms). The retrieved unclotted blood was used to inoculate directly a 50ml serum broth (Todd Hewitt/20% inactivated foetal calf  
20 serum). The culture was constantly monitored and allowed to grow to late logarithmic phase. The presence of blood in the medium interfered with OD<sub>600</sub> readings as it was being increasingly lysed with increasing growth of the bacterium, hence the requirement to constantly monitor the culture. Upon reaching late logarithmic phase/early stationary phase, the culture was transferred to a fresh 50 ml tube in order  
25 to exclude dead bacterial cells and remaining blood cells which would have sedimented at the bottom of the tube. 0.5 ml aliquots were then transferred to sterile cryovials, frozen in liquid nitrogen and stored at -70 °C. A viable count was carried out on a single standard inoculum aliquot in order to determine bacterial numbers. This was determined to be approximately 5 X10<sup>8</sup> cfu per ml.

30

#### **Intra-peritoneal Challenge and virulence testing of Group B Streptococcus standard inoculum**

To determine if the standard inoculum was suitably virulent for use in a vaccine trial, challenges were carried out using a dose range. Frozen standard inoculum strain

5 aliquots were allowed to thaw at room temperature. From viable count data the number of cfu per ml was already known for the standard inoculum. Initially, serial dilutions of the standard inoculum were made in Todd Hewitt broth and mice were challenged intra-peritoneally with doses ranging from  $1 \times 10^8$  to  $1 \times 10^4$  cfu in a 500  $\mu$ l volume of Todd Hewitt broth. The survival times of mouse groups injected with different doses of the bacterium were compared. The standard inoculum was determined to be suitably virulent and a dose of  $1 \times 10^6$  cfu was considered close to optimal for further use in vaccine trials. Further optimisation was carried out by comparing mice challenged with doses ranging between  $5 \times 10^5$  and  $5 \times 10^6$  cfu. The optimal dose was estimated to be approximately  $2.5 \times 10^6$  cfu. This represented a 100% lethal dose and was repeatedly consistent with end-points as determined by survival times being clustered within a narrow time-range. Throughout all these experiments, challenged mice were constantly monitored to clarify symptoms, stages of symptom development as well as calculating survival times.

#### 15 **Vaccine trials**

Vaccine trials in mice were accomplished by the administration of DNA to 6 week old CBA/ca mice (Harlan, UK). Mice to be vaccinated were divided into groups of six and each group were immunised with recombinant pcDNA3.1 plasmid DNA containing a specific target-gene sequence derived using the LEEP system. A total of 100  $\mu$ g of DNA in Dulbecco's PBS (Sigma) was injected intramuscularly into the tibialis anterior muscle of both hind legs. Four weeks later this procedure was repeated using the same amount of DNA. For comparison, control mice groups were included in all vaccine trials. These control groups were either not DNA-vaccinated or were immunised with non-recombinant pcDNA3.1 plasmid DNA only, using the same time course described above. Four weeks after the second immunisation, all mice groups were challenged intra-peritoneally with a lethal dose of *S. agalactiae* serotype III (strain 97/0099). The actual number of bacteria administered was determined by plating serial dilutions of the inoculum on Todd-Hewitt/5% blood agar plates. All mice were killed 3 or 4 days after infection. During the infection process, challenged mice were monitored for the development of symptoms associated with the onset of *S. agalactiae* induced-disease. Typical symptoms in an appropriate order included piloerection, an increasingly hunched posture, discharge from eyes, increased lethargy and reluctance to move which was often the result of apparent paralysis in the lower body/hind leg region. The

latter symptoms usually coincided with the development of a moribund state at which stage the mice were culled to prevent further suffering. These mice were deemed to be very close to death, and the time of culling was used to determine a survival time for statistical analysis. Where mice were found dead, a survival time was calculated by averaging the time when a particular mouse was last observed alive and the time when found dead, in order to determine a more accurate time of death.

### Interpretation of Results

A positive result was taken as any DNA sequence that was cloned and used in challenge experiments as described above and gave protection against that challenge. DNA sequences were determined to be protective;

-if that DNA sequence gave statistically significant protection (to a 95% confidence level ( $p > 0.05$ ) as determined using the Mann-Whitney U test.

-if that DNA sequence was marginal or non-significant using Mann-Whitney but showed some protective features. For example, one or more outlying mice may survive for significantly longer time periods when compared with control mice. Alternatively, the time to first death may also be prolonged when compared to counterpart mice in control groups.

It is acceptable to allow marginal or non-significant results to be considered as potential positives when it is possible that the clarity of some results may be affected by problems associated with the administration of the DNA vaccine. Indeed, much varied survival times may reflect different levels of immune response between different members of a given group.

### Results

#### Statistical analysis of survival times - LEEP DNA immunisation and GBS challenge Trial 1 (Figure 4a)

	Mean Survival Times (hours)				
	pcDNA3.1	17(ID-8)	18(ID-9)	20(ID-25)	rib
1	26.833	14.916	27.750	30.500	88.666

2	42.333	94.000 (T)	34.333	33.333	28.166
3	47.916	45.166	41.083	34.083	37.250
4	28.333	30.750	47.083	23.500	37.250
5	42.333	74.666	94.000 (T)	94.000 (T)	94.000 (T)
6	25.333	25.000	26.166	30.500	45.750
<b>Mean</b>	<b>37.549</b>	<b>51.899</b>	<b>48.849</b>	<b>43.083</b>	<b>57.066</b>
<b>sd</b>	<b>9.3943</b>	<b>32.214</b>	<b>26.257</b>	<b>28.768</b>	<b>31.556</b>
<b>p value 1</b>		<b>0.4049</b>	<b>0.4049</b>	<b>0.5000</b>	<b>0.1481</b>
<b>p value 2</b>	<b>&gt; 39.0</b>	<b>&gt; 39.0</b>	<b>&gt; 39.0</b>	<b>&gt; 39.0</b>	

(T) - terminated at conclusion of experiment but showing symptoms of infection.

5 **p value 1** refers to statistical significance when compared to pcDNA3.1 controls.

**p value 2** refers to statistical significance when compared to rib positive control.

10

All DNA vaccine's showed a pattern of protection similar to that obtained with the rib DNA vaccine, which was initially used as a positive control.

15

### **17 (ID-8)**

Mice immunised with the '17 (ID-8)' DNA vaccine did not show significantly longer survival times when compared with the unvaccinated control group. However, there are two outlying mice one of which survived the term of the experiment despite developing symptoms. The group also exhibited a much wider range of survival times reflected by a mean survival value which is approximately 14 hours higher than that demonstrated by the unvaccinated control group.

25

### **18 (ID-9)**

Mice immunised with the '18 (ID-9)' DNA vaccine did not show significantly longer survival times when compared with the unvaccinated control group. However, there is one outlying mouse which survived the term of the experiment despite developing symptoms.

### 20 (ID-25)

Mice immunised with the '20 (ID-25)' DNA vaccine did not show significantly longer survival times when compared with the unvaccinated control group. However, there was one outlying mouse which survived the term of the experiment despite developing symptoms.

### Statistical analysis of survival times - LEEP DNA immunisation and GBS challenge Trial 2 (Figure 4b)

	Mean Survival Times (hours)			
	pcDNA	UnVacc	22(ID-10)	28(ID-13)
1	45.000	27.916	44.666	72.000 (T)
2	37.333	45.083	51.416	33.000
3	37.333	37.583	40.791	36.083
4	35.291	24.583	44.666	72.000 (T)
5	24.333	37.583	36.916	49.166
6	45.000	33.166	57.833	36.083
<b>Mean</b>	<b>35.858</b>	<b>34.549</b>	<b>43.691</b>	<b>52.449</b>
<b>sd</b>	<b>7.4342</b>	<b>8.2567</b>	<b>5.3825</b>	<b>18.850</b>
<b>p value 1</b>		<b>&gt; 39.0</b>	<b>0.1137</b>	<b>0.2340</b>
<b>p value 2</b>	<b>0.4679</b>		<b>0.0323</b>	<b>0.1137</b>

(T) - terminated at conclusion of experiment but showing symptoms of infection.

p value 1 refers to statistical significance when compared to pcDNA3.1 controls.

**p value 2** refers to statistical significance when compared to unvaccinated controls.

5 There was no significant difference in the survival times exhibited by the pcDNA3.1 and unvaccinated control groups. This is confirmed by their very similar mean survival times of 35.858 hours (pcDNA3.1) and 34.166 hours (Unvaccinated).

10

### **22 (ID-10)**

Mice immunised with the '22 (ID-10)' DNA vaccine exhibited significantly longer survival times when compared with the unvaccinated control group but not when compared with the pcDNA3.1 control group. In addition, the time to first death in this group was prolonged by approximately 12 hours when compared to the pcDNA3.1 and unvaccinated control groups. The mean survival time of 43.691 hours is also considerably higher than that determined for both control groups.

20

### **28 (ID-13)**

Mice immunised with the '28 (ID-13)' DNA vaccine did not show significantly longer survival times when compared with the pcDNA3.1 and unvaccinated control groups. However there are three outlying mice, two of which survived the term of the experiment despite showing symptoms. In addition, the time to first death in this group was prolonged by approximately 9 hours when compared to the pcDNA3.1 and unvaccinated control groups. The mean survival time of 52.449 hours is also considerably higher than that determined for both control groups, as well demonstrating a wider range of survival times.

30

### **Statistical analysis of survival times - LEEP DNA immunisation and GBS challenge Trial 3 (Figure 4c)**

35



	<b>Mean Survival Times (hours)</b>				
	UnVacc.	70(ID-42)	94(ID-48)	86(ID-47)	51(ID-37)
1	27.583	25.166	34.666	32.416	43.749
2	27.583	42.666	49.500	32.416	38.333
3	24.583	34.666	27.000	42.500	50.666
4	22.250	42.666	30.500	34.500	45.166
5	35.916	30.583	30.500	34.500	69.082
6	22.250	25.166	42.666	42.500	31.166
<b>Mean</b>	<b>27.583</b>	<b>35.149</b>	<b>34.433</b>	<b>35.266</b>	<b>49.399</b>
<b>sd</b>	<b>5.1691</b>	<b>7.6444</b>	<b>8.8495</b>	<b>4.1758</b>	<b>11.846</b>
<b>p value</b>		<b>0.0628</b>	<b>0.0321</b>	<b>0.0153</b>	<b>0.0041</b>

5 **p value** refers to statistical significance when compared to unvaccinated controls.

#### **70 (ID-42)**

10 Mice immunised with the '70 (ID-42)' DNA vaccine, marginally did not  
show significantly longer survival times when compared with the  
unvaccinated control group. However, the first death in this group is  
prolonged (by approximately 3 hours ) when compared with the  
unvaccinated group. In addition, the group has a mean survival time  
15 which is approximately 8 hours longer than the unvaccinated group.

#### **94 (ID-48)**

20 Mice immunised with the '94 (ID-48)' DNA vaccine exhibited  
significantly longer survival times when compared with the unvaccinated  
control group.

#### **86 (ID-47)**

Mice immunised with the '86 (ID-47)' DNA vaccine exhibited significantly longer survival times when compared with the unvaccinated control group.

5

### 51 (ID-37)

Mice immunised with the '51 (ID-37)' DNA vaccine exhibited significantly longer survival times when compared with the unvaccinated control group.

10

### Statistical analysis of survival times - LEEP DNA immunisation and GBS challenge Trial 4 (Figure 4d)

15

	Mean Survival Times (hours)	
	UnVacc	9(ID-6)
1	32.666	35.250
2	21.666	30.958
3	23.916	69.333
4	22.999	52.333
5	25.916	44.916
6	35.916	47.083
<b>Mean</b>	<b>25.432</b>	<b>46.041</b>
<b>sd</b>	<b>4.3291</b>	<b>16.096</b>
<b>p value</b>		<b>0.0101</b>

(T) - terminated at conclusion of experiment but showing symptoms of infection.

20

**p value** refers to statistical significance when compared to unvaccinated controls

### 9 (ID-6)

Mice immunised with the '39(ID-6)' DNA vaccine showed significantly longer survival times when compared with the control group.

5

### Statistical analysis of survival times - LEEP DNA immunisation and GBS challenge Trial 6 (Figure 4e)

10

	Mean Survival Times (hours)				
	pcDNA	UnVacc	32 (ID-15)	39(ID-17)	57(40)
1	33.541	36.000	25.041	52.333	28.333
2	36.750	29.999	30.458	44.750	32.708
3	36.750	32.749	44.833	44.750	36.083
4	36.750	44.500	30.458	36.250	40.333
5	29.000	28.333	64.833	36.250	72.000 (T)
6	30.750	31.666	72.000 (T)	28.583	33.750
<b>Mean</b>	<b>34.558</b>	<b>34.316</b>	<b>39.124</b>	<b>44.016</b>	<b>38.103</b>
<b>sd</b>	<b>3.4036</b>	<b>6.3921</b>	<b>16.140</b>	<b>13.833</b>	<b>12.986</b>
<b>p value 1</b>		<b>&gt; 39.0</b>	<b>0.4043</b>	<b>0.1867</b>	<b>0.4044</b>
<b>p value 2</b>	<b>0.2862</b>		<b>0.2873</b>	<b>0.0458</b>	<b>0.2113</b>

15

(T) - terminated at conclusion of experiment but showing symptoms of infection.

**p value 1** refers to statistical significance when compared to pcDNA3.1 controls

**p value 2** refers to statistical significance when compared to unvaccinated controls.

20

There was no significant difference in the survival times exhibited by the pcDNA3.1 and unvaccinated control groups. This is confirmed by their

very similar mean survival times of 34.558 hours (pcDNA3.1) and 34.316 hours (Unvaccinated).

5     **32 (ID-15)**

Mice immunised with the '32 (ID-15)' DNA vaccine did not show significantly longer survival times when compared with the pcDNA3.1 and unvaccinated control groups. However, the '32 (ID-15)' group has two outlying mice one of which survived the term of the experiment despite showing symptoms. This group also exhibits a wide range of survival times.

15     **39 (ID-17)**

Mice immunised with the '39 (ID-17)' DNA vaccine exhibited significantly longer survival times when compared with the unvaccinated control group but was not significant when compared with the pcDNA3.1 control group. The group has a considerably higher mean survival time of 44.016 hours than that determined for either of the control groups.

25     **57 (ID-40)**

Mice immunised with the '32 (ID-15)' DNA vaccine did not show significantly longer survival times when compared with the pcDNA3.1 and unvaccinated control groups. However, the '32 (ID-15)' group has one outlying mouse which survived the term of the experiment despite showing symptoms.

30     **SURVIVAL DATA-SET B**

**Statistical analysis of survival times - LEEP DNA immunisation and GBS challenge Trial 2 (Figure 5a)**

35

	Mean Survival Times (hours)		
	pcDNA	UnVacc	13(ID-72)
1	45.000	27.916	69.166
2	37.333	45.083	36.333
3	37.333	37.583	43.916
4	35.291	24.583	32.166
5	24.333	37.583	36.333
6	45.000	33.166	43.916
Mean	35.858	34.549	43.582
sd	7.4342	8.2567	14.917
p value 1		> 39.0	0.4679
p value 2	0.4679		0.1880

**p value 1** refers to statistical significance when compared to pcDNA3.1 controls.

**p value 2** refers to statistical significance when compared to unvaccinated controls.

There was no significant difference in the survival times exhibited by the pcDNA3.1 and unvaccinated control groups. This is confirmed by their very similar mean survival times of 35.858 hours (pcDNA3.1) and 34.166 hours (Unvaccinated).

### 13 (ID-72)

Mice immunised with the '13 (ID-72)' DNA vaccine did not show significantly longer survival times when compared with the pcDNA3.1 and unvaccinated control groups. However, there is one outlying mouse which survived approximately 24 hours longer than the longest surviving mice in the pcDNA3.1 and unvaccinated control groups respectively. In addition, the time to first death in this group was prolonged when

compared to the pcDNA3.1 and unvaccinated control groups. The mean survival time of 43.582 hours is considerably higher than that determined for both control groups.

5

10

**Statistical analysis of survival times - LEEP DNA immunisation and GBS challenge Trial 3 (Figure 5b)**

	<b>Mean Survival Times (hours)</b>		
	<b>UnVacc</b>	<b>3-60(ID-65)</b>	<b>3-5(ID-66)</b>
<b>1</b>	27.583	54.416	42.916
<b>2</b>	27.583	31.000	42.916
<b>3</b>	24.583	43.000	32.874
<b>4</b>	22.250	34.916	42.916
<b>5</b>	35.916	38.958	27.333
<b>6</b>	22.250	34.916	30.916
<b>Mean</b>	<b>27.583</b>	<b>40.458</b>	<b>37.791</b>
<b>sd</b>	<b>5.1691</b>	<b>8.9959</b>	<b>7.2860</b>
<b>p value</b>		<b>0.0098</b>	<b>0.0215</b>

15

**p value** refers to statistical significance when compared to unvaccinated controls.

20

**3-60 (ID-65)**

Mice immunised with the '3-60 (ID-65)' DNA vaccine exhibited significantly longer survival times when compared with the unvaccinated control group.

5 **3-5 (ID-66)**

Mice immunised with the '3-5 (ID-66)' DNA vaccine exhibited significantly longer survival times when compared with the unvaccinated control group.

**Statistical analysis of survival times - LEEP DNA immunisation and GBS challenge Trial 4 (Figure 5c)**

	Mean Survival Times (hours)			
	UnVacc	3-40(ID-67)	3-30(ID-68)	3-38(ID-69)
1	32.666	79.750	35.500	68.583
2	21.666	35.833	28.333	29.916
3	23.916	30.500	31.208	29.916
4	22.999	22.708	98.000 (T)	31.041
5	25.916	28.583	73.500	32.166
6	35.916	40.791	32.333	29.916
<b>Mean</b>	<b>25.432</b>	<b>39.474</b>	<b>53.308</b>	<b>38.324</b>
<b>sd</b>	<b>4.3291</b>	<b>22.998</b>	<b>30.961</b>	<b>16.940</b>
<b>p value</b>		<b>0.1149</b>	<b>0.0463</b>	<b>0.1132</b>

5

(T) - terminated at conclusion of experiment but showing symptoms of infection.

p value refers to statistical significance when compared to unvaccinated controls

10

**3-40 (ID-67)**

Mice immunised with the '3-40 (ID-67)' DNA vaccine did not show significantly longer survival times when compared with the unvaccinated control group. However, there is one outlying mouse which survived approximately 43 hours longer than the longest surviving mice in the unvaccinated control group.

15

**3-30 (ID-68)**

Mice immunised with the '3-30 (ID-68)' DNA vaccine exhibited significantly longer survival times when compared with the unvaccinated control group.

20



**3-38 (ID-69)**

5 Mice immunised with the '2-19 (ID-73)' DNA vaccine did not show significantly longer survival times when compared with the unvaccinated control group. However, there was one outlying mouse which survived approximately 32 hours longer than the longest surviving mice in the unvaccinated control group. In addition, the time to first death was prolonged (by approximately 8 hours) when compared to the  
10 unvaccinated controls.

**Statistical analysis of survival times - LEEP DNA immunisation and GBS challenge Trial 5 (Figure 5d)**

	Mean Survival Times (hours)				
	UnVacc	141(ID-70)	3-20(ID-71)	2-19(ID-73)	3-6(ID-74)
1	27.833	47.500	36.166	36.166	44.666
2	45.666	52.833	44.833	49.833	36.000
3	45.666	49.333	26.750	55.833	75.416
4	34.333	46.250	36.166	68.583	36.000
5	34.333	47.500	55.916	33.333	55.916
6	45.666	36.500	44.833	30.583	36.000
<b>Mean</b>	<b>37.566</b>	<b>48.683</b>	<b>37.234</b>	<b>48.749</b>	<b>49.599</b>
<b>sd</b>	<b>7.8558</b>	<b>2.5672</b>	<b>8.4103</b>	<b>14.497</b>	<b>16.587</b>
<b>p value</b>		<b>0.0101</b>	<b>0.5000</b>	<b>0.2336</b>	<b>0.1854</b>

15

**p value** - refers to statistical significance when compared to unvaccinated controls.

**141 (ID-70)**

20 Mice immunised with the '141 (ID-70)' DNA vaccine exhibited significantly longer survival times when compared with the unvaccinated control group.

**3-20 (ID-71)**

Mice immunised with the ‘3-20 (ID-71)’ DNA vaccine did not show significantly longer survival times when compared with the unvaccinated control group. However, there is one outlying mouse which survived approximately 10 hours longer than the longest surviving mice in the unvaccinated control group.

**2-19 (ID-73)**

Mice immunised with the ‘2-19 (ID-73)’ DNA vaccine did not show significantly longer survival times when compared with the unvaccinated control group. However, there are three outlying mouse which survived approximately 4, 10 and 23 hours longer than the longest surviving mice in the unvaccinated control group. This is reflected in the higher mean survival time of 48.749 hours and a much wider range of survival times.

**3-6 (ID-74)**

Mice immunised with the ‘3-6 (ID-74)’ DNA vaccine did not show significantly longer survival times when compared with the unvaccinated control group. However, there are three outlying mouse which survived approximately 4, 10 and 23 hours longer than the longest surviving mice in the unvaccinated control group. This is reflected in the higher mean survival time of 49.599 hours and a much wider range of survival times.

**Statistical analysis of survival times - LEEP DNA immunisation and GBS challenge Trial 6 (Figure 5e)**

	Mean Survival Times (hours)			
	pcDNA3.1	UnVacc.	3-51(ID-75)	3-56 (ID-76)

1	33.541	36.000	36.333	46.583
2	36.750	29.999	30.291	29.833
3	36.750	32.749	32.000	40.166
4	36.750	44.500	52.333	46.583
5	29.000	28.333	72.000 (T)	46.583
6	30.750	31.666	40.499	---
<b>Mean</b>	<b>34.558</b>	<b>34.316</b>	<b>44.591</b>	<b>40.791</b>
<b>sd</b>	<b>3.4036</b>	<b>6.3921</b>	<b>16.615</b>	<b>7.9070</b>
<b>p value 1</b>		<b>&gt; 39.0</b>	<b>0.1876</b>	<b>0.0386</b>
<b>p value 2</b>	<b>0.2862</b>		<b>0.0867</b>	<b>0.0587</b>

(T) - terminated at conclusion of experiment but showing symptoms of infection.

5 **p value 1** refers to statistical significance when compared to pcDNA3.1 controls

**p value 2** refers to statistical significance when compared to unvaccinated controls.

10 There was no significant difference in the survival times exhibited by the pcDNA3.1 and unvaccinated control groups. This is confirmed by their very similar mean survival times of 34.558 hours (pcDNA3.1) and 34.316 hours (Unvaccinated).

15

### **3-51 (ID-75)**

Mice immunised with the '3-51 (ID-75)' DNA vaccine did not show significantly longer survival times when compared with the pcDNA3.1 control group but was relatively close to significant when compared with the unvaccinated control group. The '3-51' group has two outlying mouse one of which survived the term of the experiment despite developing symptoms. The mean survival time of 44.499 hours is considerably higher than that determined for both control groups and the group also demonstrates as a much wider range of survival times.

20

25

### 3-56 (ID-76)

5 Mice immunised with the '3-56 (ID-76)' DNA vaccine exhibited significantly longer survival times when compared with the pcDNA3.1 control group but were marginally not significant when compared with unvaccinated control group.

#### 10 **Example 3: Conservation and variability of candidate vaccine antigen genes among different isolates of Group B Streptococci**

An initial Southern blot analysis was carried out to determine cross-serotype conservation of novel Group B Streptococcal genes isolated using the LEEP system. Analysing the serotype distribution of a target gene will also determine their potential  
15 use as antigen components in a GBS vaccine. The Group B Streptococcal strains whose DNA was analysed as part of this study are listed in **APPENDIX II**.

#### **Amplification and labelling of specific target genes as DNA probes for Southern blot analysis.**

20 Oligonucleotide primers were designed for each individual gene of interest derived using the LEEP system. Primers were designed to target the whole of the gene being investigated (All primers are listed in **APPENDIX III**). Specific gene targets were amplified by PCR using Vent DNA polymerase (NEB) according to the manufacturers instructions. Typical reactions were carried out in a 100 µl volume containing 50 ng of  
25 GBS template DNA, a one tenth volume of enzyme reaction buffer, 1 µM of each primer, 250 µM of each dNTP and 2 units of Vent DNA polymerase. A typical reaction contained an initial 2 minute denaturation at 95°C, followed by 35 cycles of denaturation at 95°C for 30 seconds, annealing at the appropriate melting temperature for 30 seconds, and extension at 72°C for 1 minute (1 minute per kilobase of DNA  
30 being amplified). The annealing temperature was determined by the lower melting temperature of the two oligonucleotide primers. The reaction was concluded with a final extension period of 10 minutes at 72°C.

All PCR amplified products were extracted once with phenol chloroform (2:1:1) and once with chloroform (1:1) and ethanol precipitated. Specific DNA fragments were isolated from agarose gels using the QIAquick Gel Extraction Kit (Qiagen). For use as DNA probes, purified amplified gene DNA fragments were labelled with digoxigenin using the DIG Nucleic Acid Labelling Kit (Boehringer Mannheim) according to the manufacturer's instructions.

#### **Southern blot hybridisation analysis of Group B Streptococcal genomic DNA**

Genomic DNA had previously been isolated from all strains of Group B Streptococci which were investigated for conservation of LEEP-derived gene targets. Appropriate DNA concentrations were digested using either *Hin* DIII, *Eco* RI or *Bgl* II restriction enzymes (NEB) according to manufacturer instructions and analysed by agarose gel electrophoresis. Following agarose gel electrophoresis of DNA samples, the gel was denatured in 0.25M HCl for 20 minutes and DNA was transferred onto Hybond™ N<sup>+</sup> membrane (Amersham) by overnight capillary blotting. The method is essentially as described in Sambrook *et al.* (1989) using Whatman 3MM wicks on a platform over a reservoir of 0.4M NaOH. After transfer, the filter was washed briefly in 2x SSC and stored at 4 °C in Saran wrap (Dow chemical company).

Filters were prehybridised, hybridised with the digoxigenin labelled DNA probes and washed using conditions recommended by Boehringer Mannheim when using their DIG Nucleic Acid Detection Kit. Filters were prehybridised at 68°C for one hour in hybridisation buffer (1% w/v supplied blocking reagent, 5x SSC, 0.1% v/v N-lauryl sarcosine, 0.02% v/v sodium dodecyl sulphate[SDS]). The digoxigenin labelled DNA probe was denatured at 99.9°C for 10 minutes before being added to the hybridisation buffer. Hybridisation was allowed to proceed overnight in a rotating Hybaid tube in a Hybaid Mini-hybridisation oven. Unbound probe was removed by washing the filter twice with 2x SSC- 0.1% SDS for 5 minutes at room temperature. For increased stringency filters were then washed twice with 0.1x SSC-0.1% SDS for 15 minutes at 68°C. The DIG Nucleic Acid Detection Kit (Boehringer Mannheim) was used to immunologically detect specifically bound digoxigenin labelled DNA probes.

### Results of Southern blot analysis

All genomic digests and their corresponding Southern blots followed an identical lane order as described in Table I.

5 **Table I**

Lane	1	2	3	4	5	6	7
Strain	1 kb molecular Weight Marker	515	A909	SB35	H36B	18RS21	1954/92
Serotype		Ia	Ia	Ib	Ib	II	II

Lane	8	9	10	11	12	13	14
Strain	118/158	97/0057	BM110	BS30	M781	97/0099	3139
Serotype	II	II	III	III	III	III	IV

Lane	15	16	17	18	19	20
Strain	1169-NT	GBS 6	7271	JM9	Group A Streptococcus	<i>Streptococcus pneumoniae</i>
Serotype	V	VI	VII	VIII	—	14

10 For comparative purposes, it was decided to analyse the serotype distribution of the GBS *rib* gene, which encodes the known protective immunogen Rib. Rib has previously been shown to be present in serotype III and some strains of serotype II but not in serotypes Ia or Ib (Stalhammar-Carlemalm *et al.*, 1993). Confirmation of this pattern would not only give increased confidence in interpreting subsequent results, it would also determine if a *rib* gene homologue was present in the remaining GBS

serotypes being investigated here. Primers designed for the amplification of *rib* and its subsequent cloning into pcDNA3.1 (**Appendix I**), were used to generate a *rib* gene probe for Southern blot analysis.

5 **Southern blot analysis - *rib* (Figure 6)**

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20

10

Genomic DNA from each strain was digested completely with *Hin* DIII (NEB) and electrophoresed at 40 Volts for 6 hours in 0.8% agarose, transferred onto Hybond N<sup>+</sup> (Amersham) membrane by Southern blot and hybridised with the digoxigenin-labelled *rib* gene probe. Specifically bound DNA probe was identified using the DIG Nucleic Acid Detection Kit (Boehringer Mannheim).

15

Comment

The Southern blot analysis described in Figure 7 indicates that the *rib* gene is not conserved across all GBS serotypes. *rib* appears to be absent from all serotype Ia and Ib strains (lanes 2 to 5) and from strains 118/158 and 97/0057 of serotype II (lanes 8 and 9). However, *rib* would appear to present in strains 18RS21 and 1954/92 of serotype II (lanes 6 and 7) and in all strains of serotype III (lanes 10 to 13). This is in agreement with previously published data (Stalhammar-Carlemalm *et al.*, 1993). *rib* would also appear to be present in strains representing serotypes VII and VII (lanes 17 and 18) but was absent from strains representing serotypes IV, V and V (lanes 14 to 16) as well as the control strains (lanes 19 and 20). The *rib* gene probe did hybridise with lower intensity to genomic DNA fragments from strains representing serotypes Ia, Ib, IV, VI, VII and serotype II strains 118/158 and 97/0057. This may indicate the presence of a gene in these strains with a lower level of homology to *rib*. These hybridising DNA fragments may contain a homologue of the GBS *bca* gene encoding the Ca protein antigen which has been shown to be closely homologous to the Rib protein (Wastfelt *et al.*, 1996). If this is the case, it would be in agreement with previous work which showed all strains of serotypes Ia, Ib, II and III to be positive for one the two proteins (Stalhammar-Carlemalm *et al.*, 1993). However, the apparent

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25

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variable distribution of the *rib* gene amongst different GBS serotypes, makes it a less than ideal candidate for use in a GBS vaccine that is cross-protective against all serotypes.

5 **Southern blot analysis - 4 (ID-1) (photograph 7)**

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20

10 Genomic DNA from each strain was digested completely with *Hin* DIII (NEB) and electrophoresed at 40 Volts for 6 hours in 0.8% agarose, transferred onto Hybond N<sup>+</sup> (Amersham) membrane by Southern blot and hybridised with the digoxigenin-labelled 4 (ID-1) gene probe. Specifically bound DNA probe was identified using the DIG Nucleic Acid Detection Kit (Boehringer Mannheim).

15

Comment

The Southern blot analysis described in Figure 7 indicates that gene 4 (ID-1) is conserved across all GBS serotypes. The gene probe hybridised specifically to a *Hin* DIII-digested genomic DNA fragment of approximately 3.5 kb in DNA digests from all GBS representatives. but was absent from both the control strains (lanes 19 and 20).

20

**Southern blot analysis - 5 (ID-2) (Figure 8)**

25

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20

30 Genomic DNA from each strain was digested completely with *Eco* RI (NEB) and electrophoresed at 40 Volts for 6 hours in 0.8% agarose, transferred onto Hybond N<sup>+</sup> (Amersham) membrane by Southern blot and hybridised with the digoxigenin-labelled 5 (ID-2) gene probe. Specifically bound DNA probe was identified using the DIG Nucleic Acid Detection Kit (Boehringer Mannheim).

Comment



The Southern blot analysis described in Figure 7 indicates that gene 4 (ID-1) is conserved across all GBS serotypes. The gene probe hybridised specifically to a *Eco* RI-digested genomic DNA fragment of approximately 6.2 kb in DNA digests from all GBS representatives. but was absent from both the control strains (lanes 19 and 20).

5

#### **Southern blot analysis - 15 (ID-7) (Figure 9)**

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20

10

Genomic DNA from each strain was digested completely with *Eco* RI (NEB) and electrophoresed at 40 Volts for 6 hours in 0.8% agarose, transferred onto Hybond N<sup>+</sup> (Amersham) membrane by Southern blot and hybridised with the digoxigenin-labelled 15 (ID-7) gene probe. Specifically bound DNA probe was identified using the DIG Nucleic Acid Detection Kit (Boehringer Mannheim).

15

#### Comment

The Southern blot analysis described in Figure 7 indicates that gene 15 (ID-7) is conserved across all GBS serotypes. The gene probe hybridised specifically to a *Eco* RI-digested genomic DNA fragment of approximately 6.2 kb in DNA digests from all GBS representatives. but was absent from both the control strains (lanes 19 and 20). The gene probe hybridised specifically with *Eco* RI -digested DNA fragments ranging from approximately 3.5 kb to 5.2 kb in size.

20

25

#### **Southern blot analysis - 17 (ID-8) (Figure 10)**

#### **Figure 5**

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20

30

Genomic DNA from each strain was digested completely with *Hin* DIII (NEB) and electrophoresed at 40 Volts for 6 hours in 0.8% agarose, transferred onto Hybond N<sup>+</sup> (Amersham) membrane by Southern blot and hybridised with the digoxigenin-labelled

17 (ID-8) gene probe. Specifically bound DNA probe was identified using the DIG Nucleic Acid Detection Kit (Boehringer Mannheim).

#### Comment

5 The Southern blot analysis described in Figure 7 indicates that gene 17 (ID-8) is conserved across all GBS serotypes. The gene probe hybridised specifically to a *Hin* DIII-digested genomic DNA fragment of approximately 2.3 kb in DNA digests from all GBS representatives. but was absent from both the control strains (lanes 19 and 10 20).

#### **Southern blot analysis - 22 (ID-10) (Figure 11)**

#### **Figure 6**

15 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20

Genomic DNA from each strain was digested completely with *Bgl* II (NEB) and electrophoresed at 40 Volts for 6 hours in 0.8% agarose, transferred onto Hybond N<sup>+</sup> (Amersham) membrane by Southern blot and hybridised with the digoxigenin-labelled 20 22 (ID-10) gene probe. Specifically bound DNA probe was identified using the DIG Nucleic Acid Detection Kit (Boehringer Mannheim).

#### Comment

25 The Southern blot analysis described in Figure 7 indicates that gene 22 (ID-10) is conserved across all GBS serotypes. The gene probe hybridised specifically to a *Bgl* II-digested genomic DNA fragment of approximately 3.1 kb in DNA digests from all GBS representatives except serotype Ib strain H36B, where the gene probe hybridised specifically to a *Bgl* II-digested genomic DNA fragment. Gene 22 (ID-10) was absent from both the control strains (lanes 19 and 20).

30

#### **Conclusion**

The Southern blot analyses described here, represents a preliminary investigation into the conservation level of LEEP-derived genes amongst different GBS serotypes. Initial results indicate that the genes 4 (ID-1), 5 (ID-2), 15 (ID-7), 17(ID-8) and 22

(ID-10) are present in all GBS serotypes and thus represent potential candidate genes for use in a GBS vaccine. Similar analyses are being currently carried out for each of the genes contained in this patent.

**APPENDIX I****ID-8 (17)**

Forward Primer

5' - cgggatccgccaccatgACCACTTCTCAAGCTGTTTTAGC - 3'

Reverse Primer

5' - ttgcggccgcACGATTATCAACAAAGTTCTG - 3'**ID-9 (18)**

10 Forward Primer

5' - cggatccgccaccatgGCTACTCATATTGGAAGTTACCAGC - 3'

Reverse Primer

5' - ttgcggccgcAGGGTTTATTTGTTGAAGTGTCTTG - 3'15 **ID-10 (22)**

Forward Primer

5' - cggatccgccaccatgTATCTATATCATTACCAATGCC - 3'

Reverse Primer

5' - ttgcggccgcTTTATGTATAGAAACAGCAGTCCC - 3'

20

**ID-13 (28)**

Forward Primer

5' - cggatccgccaccatgAAAGGAAGAACAACCTATTCGTTTAG - 3'

Reverse Primer

25 5' - ttgcggccgcAAGAGCAAATTTTCGTATCTCCTC - 3'**ID-15 (32)**

Forward Primer

5' - cggatccgccaccATGATTGTTGGACACGGAATTG - 3'

30 Reverse Primer

5' - ttgcggccgcTTTTTCTTCCTCCAAAATAACACTAGC - 3'**ID-17 (39)**

Forward Primer

5' - cggatccgccaccatgGCGACTAAAGAGTTAGGTGTTAG -3'

Reverse Primer

5' - ttgcggccgcTATAGTTTTAGTTTCAACTTGTCTAGATG -3'

5 ID-25 (20)

Forward Primer

5' - cgggatccaccatgTATACGAGTTTACAACCAAATCATG -3'

Reverse Primer

5' - ttgcggccgcGTCAGCTCGTACTGTTTTTTTAGC -3'

10

ID-37 (51)

Forward Primer

5' - cggatccgccaccatgTGTCAAATGAATAGTGAACATAAAAAG -3'

Reverse Primer

15

5' - ttgcggccgcCTCAAATAATTTACCTCCAATTCG -3'

ID-40 (51)

Forward Primer

5' - cggatccgccaccatgGCTCCATTCGAATTTAAAGATTC -3'

20

Reverse Primer

5' - ttgcggccgcTGATTTACCAGTTTGGGAAGAGTTC -3'

ID-42 (70)

Forward Primer

25

5' - cggatccgccaccATGAATACTATTTATAATACATTGAGAACAG -3'

Reverse Primer

5' - ttgcggccgcTTCTTTGTTCCAACCTTTCTGG -3'

ID-47 (86)

30

Forward Primer

5' - cggatccgccaccATGATAGAGTGGATTCAAACACATTTAC -3'

Reverse Primer

5' - ttgcggccgcTTTATGACTCAAGCGACGTGTTA -3'

ID-48 94

Forward Primer

5' - cggatccgccaccATGGAGTTAGTAATTAGAGATATTCGTAAG

Reverse Primer

5' - ttgcggccgcCTTGTCATATTCATCTCCCTTCAACID-67 (3-40)

Forward Primer

5' - cggatccgccaccatgGCTAGTTTTGTCATGAATCATAATGAC -3'

10 Reverse Primer

5' - ttgcggccgcGTTATTTGCTCGTTGTTTAGCTAAATC -3'ID-68 (3-30)

Forward Primer

15 5' - cggatccgccaccatgGCTCTTAGTTTTTTTTATGGTTTCAGTTCAAGC -3'

Reverse Primer

5' - ttgcggccgcGAAGGCACCGCCACCTCC -3'ID-69 (3-38)

20 Forward Primer

5' - cggatccgccaccatgGGTGAAACCCAAGATACCAATCAAGC -3'

Reverse Primer

5' - ttgcggccgcAACACCTGGTGGGCGTTTGG -3'25 ID-70 (141)

Forward Primer

5' - cggatccgccaccATGGCTGGGAATCGTAATAACG -3'

Reverse Primer

30 5' - ttgcggccgcAGCCGTCTCTAAAACAGGCTTG -3'ID-71 (3-20)

Forward Primer

5' - cggatccgccaccatgCTTCCAACGCAGCCGCAAAC -3'

Reverse Primer

5' - ttgcgccgcATTTAGTGTTATTTCTCCTGTTGCATAATCC -3'

ID-72 (13)

Forward Primer

5' - cgggatccaccatgTACACGCATATTGTTGAAAAAAG -3'

Reverse Primer

5' - ttgcgccgcAAATAATTTCTTTTGGTTGTTTG -3'

ID-73 (2-19)

10 Forward Primer

5' - cggatccgccaccatgAGTAATCAAGAAGTTTCAGCAAGC -3'

Reverse Primer

5' - ttgcgccgcCCATTGTGGAATATCAGCTGAAG -3'

15 ID-74 (3-6)

Forward Primer

5' - cggatccgccaccatgGTGCAGGCAGTGGTACCGCT -3'

Reverse Primer

20 5' - ttgcgccgcGCGCATTGTAACAAATTCCTCAG -3'

ID-75 (3-51)

Forward Primer

5' - cgggatccaccatgGCTGCCGAGAAGGATAAAG -3'

25 Reverse Primer

5' - ttgcgccgcATTATTTAGCTGCTTTTTTAATGG -3'

ID-76 (3-56)

Forward Primer

30 5' - cgggatccaccatgTGTCAGGTTGTTTATGCAAGTTTTC -3'

Reverse Primer

5' - ttgcgccgcTTTACTAATTGATAAAGAGCAACTTCG -3'

*rib* (control)

Forward primer

5' - ggggtaccggccaccATGGCTGAAGTAATTCAGGAAGT -3'

Reverse primer

5' - cggaattccgTTAATCCTCTTTTTTTCTTAGAACAGAT



**APPENDIX II**

Listed below are the details (serotype and strain designation) of Group B Streptococcus strains whose DNA was analysed for gene conservation

5

	<b>SEROTYPE</b>	<b>STRAIN</b>
	Ia	515
	Ia	A909
10	Ib	SB35
	Ib	H36B
	II	18RS21
	II	1954/92
	II	118/158
15	II	97/0057
	III	BM110
	III	BS30
	III	M781
	III	97/0099
20	IV	3139
	V	1169/NT
	VI	GBS VI
	VII	7271
	VIII	JM9

25

A group A Streptococcal strain (serotype M1, strain NCTC8198) and *Streptococcus pneumoniae* (serotype 14) were also included in the analysis for control purposes.

**APPENDIX III**ID-1 (4)

forward primer

5 5' - atggaaaaaataacttgaaaaaattac -3'

reverse primer

5' - ctattingtttagcgatgtctttatc -3'

ID-2 (5)

10 forward primer

5' - atgtcaaaacaaaaagtaacggcaac -3'

reverse primer

5' - ttattatggccaataaccataagtaattg

15 ID-6 (9)

forward primer

5' - atgaaaaagttttttctcatggctatg -3'

reverse primer

5' - ttacttcaactgttgatagagcaacttc - 3'

20

ID-7 (15)

forward primer

5' - ttgtcaatttataggtttagaacttgg -3'

reverse primer

25 5' - ttaatttcattgcgtctcaaacc -3'

ID-8 (17)

forward primer

5' - atgacaaaaaacttattattgctatattag -3'

30 reverse primer

5' - ttaacgattatcaacaaagtctgtac -3'

ID-10 (22)

forward primer

5' - atgatacgccagtttttaagagaa -3'

reverse primer

5' - ttatttatgtatagaaacagcagtc -3'

## 5 References

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- Wastfelt, M., Stalhammar-Carlemalm, M., (1996) Identification of a family of  
Streptococcal surface proteins with extremely repetitive structure. *J. Biol. Chem.* **271**:  
18892-18897.
- 30 Zhang, D., Yang, X., Berry, J. Shen, C., McClarty, G. and Brunham, R.C. (1997) DNA vaccination with the major outer-membrane protein genes induces acquired immunity to *Chlamydia trachomatis* (mouse pneumonitis) infection. *Infection and Immunity*, **176**, 1035-40.

## Claims:

1. A *Group B Streptococcus* protein having a sequence selected from those described in fig 1, or fragments or derivatives thereof.  
5
2. A *Group B Streptococcus* polypeptide or peptide having a sequence selected from those described in fig 2, or fragments or derivatives thereof.
3. Derivatives or variants of the proteins, polypeptides, and peptides as claimed in claims 1 and 2 which show at least 50% identity to those proteins, polypeptides and peptides claimed in claims 1 and 2.  
10
4. A nucleic molecule comprising or consisting of a sequence which is:  
15
  - (i) any of the DNA sequences set out in figure 1 and figure 2 herein or their RNA equivalents;
  - (ii) a sequence which is complementary to any of the sequences of (i);
  - (iii) a sequence which codes for the same protein or polypeptide, as those sequences of (i) or (ii);  
20
  - (iv) a sequence which shows substantial identity with any of those of (i), (ii) and (iii); or
  - (v) a sequence which codes for a derivative, or fragment of a nucleic acid molecule shown in figure 1 or figure 2.
- 25 5. A vector comprising DNA encoding for the expression of any one or more proteins, polypeptides, peptides, fragments or derivatives thereof, as claimed in claims 1 to 3.
- 30 6. A vector as claimed in claim 5 further comprising DNA encoding any one or more of the following: promoters, enhancers, signal sequences, leader sequences,

translation start and stop signals, DNA stability controlling regions, or a fusion partner.

- 5 7. The use of a vector as claimed in claims 5 and 6 in the transformation or transfection of a prokaryotic or eukaryotic host.
8. A host cell suitable for the transformation of vector as claimed in claims 5 and 6.
- 10 9. An antibody, an affibody, or a derivative thereof which binds to one or more of the proteins, polypeptides, peptides, fragments or derivatives thereof, as claimed in any one of claims 1 to 3.
- 15 10. An immunogenic composition comprising one or more of the proteins, polypeptides, peptides, fragments or derivatives thereof, or nucleic acid sequences as claimed in any one or more of claims 1-3 and claim 4.
11. An immunogenic composition as claimed in claim 10 which is a vaccine.
- 20 12. Use of an immunogenic composition as a claimed in claim 10 in the preparation of a medicament for the treatment or prophylaxis of *Group B Streptococcus* infection.
- 25 13. A method of detection of *Group B Streptococcus* which comprises the step of bringing into contact a sample to be tested with at least one antibody, affibody, or a derivative thereof, as described herein.
- 30 14. A method of detection of *Group B Streptococcus* which comprises the step of bringing into contact a sample to be tested with at least one protein, polypeptide, peptide, fragments or derivatives as described herein.

15. A method of detection of *Group B Streptococcus* which comprises the step of bringing into contact a sample to be tested with at least one nucleic acid molecule as described herein.

5

16. A kit for the detection of *Group B Streptococcus* comprising at least one antibody, affibody, or derivatives thereof as claimed in claim 9.

10

17. A kit for the detection of *Group B Streptococcus* comprising at least one *Group B Streptococcus* protein, polypeptide, peptide, fragment or derivative thereof as claimed in claims 1 to 3.

18. A kit for the detection of *Group B Streptococcus* comprising at least one nucleic acid molecule as claimed in claim 4.

15

19. A method of screening for DNA encoding bacterial cell envelope associated or surface antigens in gram positive bacteria comprising the steps of:

- combining a reporter vector including the nucleotide sequence encoding the mature form of the staphylococcus nuclease gene and an upstream promoter region with DNA from a gram positive bacteria.
- transforming the resultant vector into *Lactococcus lactis* cells.
- assaying for the secretion of staphylococcus nuclease protein in the transformed cells.

20

20. A method as claimed in claim 19 wherein the reporter vector is one of the pTREP1-*nuc* vectors shown in figure 4.

25

21. A method as claimed in claim 19 or claim 20 wherein the gram positive bacteria is *Group B Streptococcus*, *Streptococcus Pneumoniae*, *Staphylococcus aureus* or pathogenic group A streptococci.

30

22. A vector as shown in figure 4 for use in screening for DNA encoding bacterial cell envelope associated or secreted antigens in gram positive bacteria.
- 5 23. A method of determining whether a protein, polypeptide, peptide, fragment or derivative thereof as claimed in claims 1 to 3 represents a potential anti-microbial target which comprises inactivating said protein and determining whether *Group B Streptococcus* is still viable.

ID-1

## FIG. 1

Clone 4

ATGGAAAAAATACTTGGAAAAAATTACTTGTTAGTACTGCTGCTCTTTCAGTAGT  
TGCAGGAGGAGCAATTGCTGCTACTCACTCTAACTCAGTTGATGCTGCTTCAAAAA  
AACTATCAAACCTTTGGGTCCCAACAGATTCAAAGCGTCTTATAAAGCAATTGTT  
AAAAAATTCGAGAAGGAAAAACAAGGCGTACTGTAAAAATGATTGAGTCTAATG  
ACTCCAAAGCTCAAGAAAACGTAAAAAAGACCCAAGCAAGGCAGCCGATGTATT  
CTCACTTCCACATGACCAACTTGGTCAATTAGTAGAATCTGGTGTATCCAAGAAA  
TTCCAGAGCAATACTCAAAGAAATTGCTAAAAACGACACTAAACAATCACTTAC  
TGGTGCACAATATAAAGGGAAAACCTTATGCATTCCCATTTGGTATTGAATCTCAAG  
TTCTTTATTATAATAAAACAAAGTTAACTGCTGACGACGTTAAATCATAACGAAACA  
ATTACAAGCAAAGGGAAATTTCGGTCAACAGCTTAAAGCAGCTAACTCATATGTAA  
CAGGTCCTCTTTTCCTTCTGTAGGCGACACTTTATTTGGTAAATCTGGTGAAGATG  
CTAAAGGCACTAACTGGGGTAATGAAGCAGGTGTTTCTGTCTTAAATGGATTGCA  
GATCAAAGAAAAATGATGGTTTTGTCAACTTGACAGCTGAAAATACAATGTCTAA  
ATTTGGCGATGGTTCTGTTTCATGCTTTTGAAAGTGGACCATGGGATTACGACGCTG  
CTAAAAAAGCTGTTCGGTGAAGATAAAATCGGTGTTGCTGTTACCCAACAATGAAA  
ATCGGTGACAAAGAAGTTCAACAAAAAGCATTCTTGGGCGTTAACTTTATGCCGT  
TAACCAAGCACCTGCTGGTTCAAACACTAAACGAATCTCAGCTAGCTACAACTCG  
CTGCATATCTAACTAATGCTGAAAGTCAAAAAATTCAATTCGAAAAACGTCATATC  
GTTCTGCTAACTCATCAATTCATCTTCTGATAGCGTCCAAAAAGATGAACTTGC  
AAAAGCAGTTATCGAAATGGGTAGCTCAGATAAATATACAACGGTTATGCCTAAG  
TTGAGTCAAATGTCAACATTCTGGACAGAAAGTGCTGCTATTCTTAGCGATACTTA  
CAGTGGTAAATCAAATCTAGCGATTACCTTAAACGTCTAAAACAATTCGATAAAG  
ACATCGCTAAAACAAAATAG

MEKNTWKLLVSTAALSVVAGGAIAATHSNSVDAASKKTIKLWVPTDSKASYKAIVK  
KFEKENKGVTVKMIENSNSKAQENVKKDPSKAADVFLPHDQLGQLVESGVIQEIQ  
YSKEIAKNDTKQSLTGAQYK GKTYAFPFGIESQVLYYNKTKLTADDVKS YETITSK GK  
FGQQLKAANSYVTGPLFLSVGDTLFGKSGEDAKGTNWGNEAGVSVLKWIADQKKN  
GFVNLT AENTMSKFGDGSVHAFESGPWDYDAKKA VGEDKIGVAVYPTMKIGDKEV  
QQKAFLGVKLYAVNQAPAGSN TKRISASYKLAAYLTNAESQKIQFEKRHIVPANSSIQS  
SDSVQKDELAKAVIEMGSSDKYTTVMPKLSQMSTFWTESAAILSDTYS G KIKSSDY  
LKRLKQFDKDIATKZ

ID-2

Clone 5



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ATGTCAAACAAAAAGTAACGGCAACTTTGTTGTTATCCACTTTAGTCTTATCGCT  
 ATCATCACCTTTAGTGACCTTAGCAGAAACTATTAATCCAGAAACAAGCCTGACAA  
 TGGCAACAGCATCAACAGAAAGTTCTTCTGAAGCAGAGAAACAGGAAAAAACACA  
 ACCTACAGATTCAGAAACTGCTTCACCTTCAGCCGAAGGAAGTATCTCAACAGAA  
 AAAACAGAGATTGGTACGACAGAGACATCATCAAGCAATGAATCATCATCAAGTT  
 CATCACATCAATCTTCTTCCAACGAAGATGCTAAAACATCTGATTCTGCTTCAACA  
 GCATCTACTCCTAGCACTAATACTACAAACAGTAGTCAAGCAGACAGTAAGCCAG  
 GTC AATCAACAAAGACTGAATTA AACCTGAGCCTACCTTACCATTAGTAGAGCCT  
 AAAATAACTCCCGCTCCGTCTCAGATAGAAAGTGTT CAGACAAATCAGAATGCTTC  
 TGTTCCCTGCTTTATCCTTTGATGATAACTTATTATCAACACCGATTCACCAGTGAC  
 AGCAACGCCATTCTACGTAGAACACTGGTCTGGTCAGGATGCCTACTCTCACTATT  
 TATTGTCACATCGTTACGGTATCAAAGCTGAACAATTAGATGGGTA CTTAAAATCT  
 TTAGGGATTCAATATGATTCTAATCGTATCAATGGTGCTAAGTTATTACAATGGGA  
 AAAAGATAGTGGTTTAGATGTCCGTGCTATTGTAGCTATTGCTGTCCTTGAAAGTTC  
 ATTGGGAACTCAAGGAGTGGCTAAAATGCCAGGTGCTAATATGTTTGGTTATGGTG  
 CCTTTGATCATGACTCTAGCCATGCTAGTGCTTATAATGATGAAGAAGCAATTATG  
 TTGTTGACAAAAAATACAATTATTA AAAACAACA ACTCTAGCTTTGAAATCCAAGA  
 TTTGAAAGCACAGAAATTATCTTCTGGACA ACTTAATACAGTTACTGAGGGTGGTG  
 TTTATTATACAGATAACTCTGGA ACTGGTAAACGTCGTGCC CAGATTATGGAAGAT  
 TTAGACCGCTGGATTGATCAACATGGAGGGACACCAGAAATTCCTGCTGCCTTGAA  
 AGCTTTATCGACAGCAAGTTTAGCAGATTACCAAGTGGTTTTAGCTTATCAACAG  
 CGGTTAACACAGCTAGCTATATTGCATCAACTTATCCATGGGGTGAATGTACATGG  
 TATGTCTTTAACC GCGCTAAAGAGTTAGGTTATACATTTGATCCATTTATGGGTAAT  
 GGTGGAGATTGGCAACATAAGGCTGGCTTTGAAACAACACATTCACCAAAGTAG  
 GCTATGCTGTATCATTTTCACCAGGACAAGCTGGTGCTGATGGCACTTACGGTCAC  
 GTAGCTATTGTTGAAGAAGTTAAAAAAGATGGTTCAGTTCTCATTT CAGAATCTAA  
 TGCAATGGGACGTGGTATTGTCTCTTACCGTACTTTTAGTT CAGCACAAGCTGCAC  
 AATTA ACTTATGGTATTGGCCATAAATAA

MSKQKVTATLLLSTLVLSLSSPLVTLAETINPETSMTASTESSSEAEKQEKTPQPTDS  
 ETASPSAEGSISTEKTEIGTTETSSSNESSSSSSHQSSSNEDAKTSDSASTASTPSTNTNS  
 SQADSKPGQSTKTELKPEPTLPLVEPKITPAPSQIESVQTNQNASVPALSFDDNLLSTPIS  
 PVTATPFYVEHWGQDAYSHYLLSHRYGIKAEQLDGYLKSLGIQYDSNRINGAKLLQ  
 WEKDSGLDVRAIVAIAVLESSLGTQGVAKMPGANMFGYGAFDHDSSHASAYNDEEAI  
 MLLTKNTIKNNNSSFEIQDLKAQKLSSGQLNTVTEGGVYYTDNSGTGKRRAQIMEDL  
 DRWIDQHGGTPEIPAALKALSTASLADLPSGFSLSAVNTASYIASTYPWGECTWYVF  
 NRAKELGYTFDFPMGNGGDWQHKAGFETTHSPKVGYAVSFSPGQAGADGTYGHVAI  
 VEEVKKDGSVLISESNAMGRGIVSYRTFSSAQAAQLTYGIGHKZ

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FIG. 1 CONT'D

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ID-3

Clone 6

GTGCATATGTTACAAAACATTGGACAAACAGGCATTCAAGCAACTCGAATTGCTTT  
AGGTTGTATGAGAATGAGTGACTTGAAAGGAAAACAAGCTGAAGAAGTAGTTGGA  
ACAGCATTAGATTTGGGTATTATAAATAATAAAGTGCAAGAAAGTGTCTCTGGCGT  
CAAAGTGACTIONAATCATTGTGTTATCAAGAACAAGAAATTGCTTCTTTTCAAGAGA  
TTAATCAGATGACTTTCGTGAAGAACATGCGGACCATGACTTATGATGTCATGTTT  
GATCCTTTAGTTCTTCTTTTTATAGGTGCCTCCTACGTATTAACATTGGCTATGGGA  
GCTTTTATGATTTCAAAGGTCAAGTTACTGTTGGTGACTTGGTAACATTTGTGACG  
TATTTAGATATGTTGGTATGGCCCTTGATGGCGATTGGTTTCTTGTTCAATATGGTA  
CAGCGTGGTAGTGTCTTATAACCGTATTAATAGTCTACTTGAGCAAGAATCGGA  
TATAACTGATCCTTTAAATCCTATCAAACCTGTTGTCAATGGAACATTAAGATA  
TGATATTGATTTCTTTAGATACGACAATGAGGAAACCTTAGCCGATATTCATTTAC  
CTTAGAAAAAGGTCAAACCTTAGGTTTGGTAGGTCAAACGGGATCAGGGAAGACA  
AGTCTTATTAAGTTATTGCTACGTGAACATGATGTGACTCAGGGGAAAATTACTTT  
AAATAAACATGATATACGTGATTATCGATTGTCTGAGTTACGTCAACTAATCGGTT  
ATGTTCTCAAGATCAGTTTTTATTTGCTACCAGTATTTTAGAAAATGTTTCGCTTTG  
GAAATCCAACCTCTATCTATCAATGCTGTCAAAGAAGCAACTAAATTGGCACATGTT  
TACGATGACATTGAACAGATGCCAGCAGGATTTGAGACTCTAATTGGAGAAAAAG  
GAGTCTCATTATCTGGTGGACAAAAACAAGGATTGCGATGAGTCGTGCCATGATT  
TTAGATCCAGATATTCTTATTTTGGATGATTCTCTATCAGCAGTGGACGCTAAAACG  
GAACATGCTATTGTTGAGAATCTTAAAACGAATCGTCAAGGGAAATCGACTATTA  
TTTCAGCACATCGTTTATCAGCTGTTGTGCACGCAGACCTTATCTTAGTTATGCGAG  
ACGGCAGAGTCATTGAGCGAGGTCAACATCAAGAGTTGCTAAATAAAGGTGGTTG  
GTATGCTGAAACGTATGCCTCACAGCAATTAGAAATGGAGGAAGCATTGATGAA  
GTCTAA

MHMLQNIQGTGIQATRIALGCMRMSDLKKGKQAEVVGTALDLGIINNKVQESVSGVK  
VTKSLCYQEQEIASFQEINQMTFVKNMRTMTYDVMFDPLVLLFIGASYVLTAMGAF  
MISKGQVTVGD LVTFV TYLDMLVWPLMAIGFLFNMVQRGSVSYNRINSLEQESDITD  
PLNPIKPVVNGTLRYDIDFFRYDNEETLADIHFTLEKGQTLGLVGQTGSGKTSLIKLLR  
EHDVTQGKITLNKHDIRDYRLSELRQLIGYVPQDQFLFATSILENVRFGNPTLSINAVKE  
ATKLAHVYDDIEQMPAGFETLIGEKGVSLSGGQKQRIAMSRAMILDPDILILDDSLSAV  
DAKTEHAIVENLKTNRQKGSTIISAHRLSAVVHADLILVMRDGRVIERGQHQELLNKG  
GWYAETYASQQLEMEEAFDEVZ

ID-4

Clone 6b

TTGATGAAGTCTAATCAATGGCAAGTCTTTAAGAGATTAATCTCCTATTTACGCCCT  
TATAAATGGTTTACAGTATTAGCTCTATCTCTTATTGTTGACGACTGTTGTAAA

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FIG. 1 CONT'D

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AATATTATTCCTTTAATTGCTTCACATTTTATTGATCACTATCTGACAAATGTTAAT  
 CAAACAGCAGTTCTTATTTTAGTGGGATATTATTCAATGTATGTCTTGCAGACCTTA  
 ATTCAATATTTTGGGAATCTCTTTTTTTCGCGGTGTTTCTTATAGTATTGTTAGAGAT  
 ATTCGTAGAGATGCTTTTGCTAATATGGAAAGGCTAGGCATGTCTTATTTTGATAG  
 GACACCGGCAGGATCTATTGTGTCACGTATTACTAATGATACTGAAGCAATATCTG  
 ATATGTTTTTCGGGTATTTTATCAAGTTTTATCTCGGCGATATTTATTTTACAGTTAC  
 TCTGTACACTATGTTGATGCTAGACATTAACACTAACAGGACTCGTCGCTCTTTTGGT  
 ACCTGTTATCTTTATATTAGTGAATGTCTATCGGAAAAAATCAGTCACTGTCATTGC  
 TAAAACGAGAAGTTTACTTAGTGATATCAACAGTAAATTATCAGGAAGTATTGAAG  
 GAATTCGCATTGTACAGGCTTTTGGTCAAGAAGAGCGCTTGAAGACTGAATTTGAG  
 GAAATTAACAAAGAGCATGTTGTGTATGCCAATCGTTCTATGGCTCTTGATAGTCT  
 CTTCTTAAGACCGGCGATGTCTCTTTTAAACTCCTAGCATATGCTGTTCTTATGTC  
 TTATTTTGGATTTACAGGAGTTAAAGGAGGTCTTACGGCAGGATTAATGTATGCTT  
 TTATTCAGTACGTTAATCGTCTATTTGACCCTTTAATTGAAGTAACGCAAAATTTT  
 CAACCTTACAAACATCAATGGTATCAGCAGGGCGTGTGTTTGATCTGATTGAT  
 GAAACAGGTTTTGAACCAAGCCAAAAAATACAGAAGCT

MKSNQWQVFKRLISYLRPYKWFTVLALSLLLLTTVVKNIIPLIASHFIDHYLTNVNQTA  
 VLILVGYYSMYVLQTLIQYFGNLFFARVSYIVRDIRRDAFANMERLGMYSYFDRTPAG  
 SIVSRITNDTEAISDMFSGILSSFISAIFIFTVTLYTMLMLDIKLTGLVALLPVIFILVNVY  
 RKKSVTVIKTRSLSDINSKLSGSIEGIRIVQAFGQEERLKTEFEEINKEHVVYANRSM  
 ALDSLFLRPAMSLKLLAYAVLMSYFGFTGVKGGTAGLMYAFIQYVNRFLFDPLIEVT  
 QNFSTLQTSMVSAGRVFDLIDETGFEPKQNTA

ID-5

Clone 7

ATGAAAAGAAAAGACTTATTTGGTGATAAACAACACTCAATACACGAT  
 TAGAAAGTTAAGTGTGGAGTAGCTTCAGTTGCAACAGGGGTATGTA  
 TTTTCTTCATAGTCCACAGGTATTTGCTGAAGAAGTAAGTGTTCCTC  
 CTGCAACTACAGCGATTGCAAAGTCGAATATTAATCAGGTTGACAAC  
 CGGCAATCTACTAATTTAAAAGATGACATAAACTCAAACCTGAGAC  
 GGTGTGACACCCTCAGATATGCCGGATACCAAGCAATTAGTATCAG  
 ATGAAACTGACACTCAAAAAGGAGTGACAGAGCCGGATAAAGGCGAC  
 AAGCCTGCTTGAAGAAAATAAAGGTCTGTTTCAGATAAAAATACT  
 TAGATTTAAAAGTGGCACCATCTACATTGCAAAATACTCCCGACAAA  
 ACTTCTCAAGCTATAGGTGCTCCAAGTCCGACCTTGAAGTTGCTAAT  
 CAAGCTCCACAGATTGAAAATGGTTACTTTAGGTTACATCTTAAAGA  
 ATTGCCTCAAGGTCATCCTGTAGAAAGCACTGGGCTTTGGATATGGG  
 GAGATGTTGATCAACCGTCTAGTAATTGGCCAAATGGTGCTATCCCT  
 ATGACTAATGCTAAGAAAGATGATTACGGTTATTATGTTGATTTTAA  
 ATTATCTGAAAAACAACGAAAACAATATCTTTTTTAATTAATAACA  
 AAGCAGGAACAAATTTAAGCGGCGATCATCATATTCCATTATTACGA

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FIG. 1 CONT'D

CCTGAGATGAACCAAGTTTGGATTGATGAAAAGTACGGTATACATAC  
TTATCAGCCCCCTCAAAGAAGGGTATGTCCGTATTA ACTATTTGAGTTC  
ATCTGGTAACTATGACCACTTATCAGCATGGCTCTTTAAAGATGTTGC  
AACCCCTCAACA ACTTGGCCAGATGGTAGTAATTTTGTGAATCAAG  
GACTATATGGAAGGTATATTGATGTACCACTGAAA ACTAATGCCAAA  
GAGATTGGTTTTCTAATCTTAGATGAAAGTAAGACAGGAGATGCAGT  
GAAAGTTCAACCCAACGACTATGTTTTTAGAGATTTAGCTAACCAT  
ACCAAATTTTTGTAAAAGATAAGGATCCAAAGGTTTATAATAATCCT  
TATTACATTGATCAAGTGCAGCTAAAGGATGCTCAACAACTGATTT  
AACAAAGTATTCAAGCAAGTTTTACA ACTCTAGATGGGGTAGATAAAA  
CTGAAATTTTAAAAGAATTGAAAGTGACAGATAAAAATCAAATGCT  
ATACAAATTTCTGATATCACTCTCGATACTAGTAAATCTCTTTAATA  
ATCAAAGGCGACTTTAATCCTAAACAAGGTCATTTCAATATATCTTAT  
AATGGTAAACAATGTCACGACAAGGCAATCTTGGGAATTTAAAGACCA  
ACTTTATGCTTATAGTGGAAATTTAGGTGCAGTTCTCAATCAAGATGG  
TTCAAAGTTGAAGCCAGCCTCTGGTCACCGAGTGCTGATAGTGCA  
CTATGATTATTTATGACAAAGATAATCAAACAGGGTGTAGCGACT  
ACCCCTTGTGAAAAATAATAAGGTGTTTGGCAGACGATACTTGA  
TACTAAATTAGGTATTA AAAACTATACTGGTACTATTATCTTTACGA  
AATAAAAAGAGGTAAGGATAAGGTTAAGATTTTAGATCCTTATGCAA  
AGTCATTAGCAGAGTGGGATAGTAATACTGTTAATGACGATATAAAA  
ACGGCTAAAGCAGCTTTTGTAATCCAAGTCAACTTGGACCTAAAAA  
TTTAAGTTTTGCTAAAATTGCTAATTTTAAAGGAAAACAAGATGCTGT  
TATATACGAAGCACATGTAAGAGACTTCACTTCTGATCAATCTTTGG  
ACGGAAAATTA AAAAATCAACTTGGTACCTTTGCAGCCTTTTCAGAG  
AACTAGATTATTTACAGAAATTAGGAGTTACACACATTCAGCTTTT  
ACCGGTATTGAGTTATTTTTATGTTAATGAAATGGATAAGTCACGCTC  
AACAGCTTACACTTCCTCAGACAATAATTACAATTGGGGCTATGACC  
CACAGAGCTATTTTGCTCTTTCTGGAATGTATTCAGAGAAACCAAAA  
GATCCATCAGCACGTATCGCCGAATTA AAAACAATTAATACATGATAT  
TCATAAACGTGGCATGGGGGTTATACTTGATGTCGTCTATAATCACA  
CTGCAAAA ACTTATCTCTTTGAGGATATAGAACCTAATTATTACT  
TTATGAATGAAGATGGTTCACCAAGAGAAAGTTTTGGAGGGGGACGT  
TTAGGAACCACTCATGCAATGAGTCGTCTGTTTTGGTTGATTCCATT  
AAATATCTTACAAGTGAATTTAAAGTTGATGGTTTCCGTTTTGATATG  
ATGGGAGATCATGATGCGGCTGCGATTGAATTAGCTTATAAAGAAGC  
TAAAGCTATTAATCCTAATATGATTATGATTGGTGAGGGCTGGAGAA  
CATTCCAAGGCGATCAAGGTAAGCCGGTTAAACCAGCTGACCAAGAT  
TGGATGAAGTCAACCGATACAGTTGGCGTCTTTTCAGATGATATTCGT  
AATAGCTTGAAATCTGGTTTTCCAATGAAGGTA CTCCAGCTTTTCATC  
ACAGGTGGCCCAATCTTTACAAGGATTTTTTAAAATATCAAAGC  
ACAACCTGGGAATTTGAAGCAGATTCGCCAGGAGATGTGGTGCAGT  
ATATTGCTGCACATGATAACCTTACCTTGCATGATGTGATTGCAAAAT  
CAATTAATAAAGACCCTAAGGTAGCTGAAGAAGATATTCATAGACGT

FIG. 1 CONT'D

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CTGCGTTTAGGAAATGTAATGATTTTAAACATCTCAAGGGACAGCATT  
CATTCAATTCTGGTCAAGAGTATGGTCGTACGAAGCGTTTACTTAACCC  
TGATTACATGACAAAAGTTTCAGATGACAAATTGCCTAATAAAGCAA  
CACTTATTGAAGCTGTAAAGAATACCCATATTTTATTCATGATTCAT  
ATGATTCTTCAGATGCCATTAATCATTTTGGATTGGGCAGCAGCCACAG  
ATAATAACAAACACCCAATTTCAACGAAAACACAGGCCTATACAGCA  
GGTTTAATCACATTAAGGCGTTCAACAGATGCTTTCCGGAAATTGAG  
CAAAGCAGAAATTGATCGTGAGGTTAGCTTGATTACAGAGGTAGGTC  
AAGGTGATATTAAGAAAAAGATTTGGTTATTGCTTACCAAACAATA  
GATTCTAAAGGCGATATTTACGCAGTATTTGTTAATGCTGATAGTAA  
AGCTAGAAACGTTTTACTAGGTGAAAAATATAAACACCTTTTAAAAG  
GGCAAGTAATTGTTGATGCTGATCAAGCGGGGATTAACCAATCTCA  
ACTCCTAGAGGTGTTCAATTTTAAAAAGATAGTTTGCTGATTGATCCA  
TTAACAGCAATTGTGATTAAGTTGGCAAAGTTGCTCCTAGCCCTAA  
GGAGGAATTGCAAGCAGATTATCCCAAACACAATCTTTCAAGGGAT  
CTAAAACGGTAGAAAAAGTAAATAGAATAGCTAATAAGACCTCAAT  
AACTCCTGTAGTTTCTAATAAGACCGATTCATATCTGACAAATGAAG  
CTAATTTGCCAAAAACTGGAGATAAGTCATCAAAAATACTAAGTGTA  
GTAGGAATAAGCATTCTAGCAAGTCTACTTGCTCTACTAGGTCTCTCT  
TTAAAGAGGAATCGCACTTAA

MKRKDLFGDKQTQYTIRKLSVGVASVATGVCIFLHSPQVFAEEVSVSPA  
TTAIAKSNNINQVDNRQSTNLKDDINSNSETVVTPSDMPDTKQLVSDTDT  
QKGVTEPDKATSLLEENKGPVSDKNTLDLKVAPSTLQNTDPKTSQAIGA  
PSPTLKVANQAPQIENGYFRLHLKELPQGHVESTGLWIWGDVDQPSSN  
WPNGAIPMTNAKKDDYGYVDFKLSEKQRKQISFLINNKAGTNLSGDH  
HIPLLRPEMNQVWIDEKYGIHTYQPLKEGYVRINYLSSSGNYDHLAWL  
FKDVA TPSTT WPDGSNFVNQGLYGRYIDVPLKTNAKEIGFLILDESKTGD  
AVKVQPNDYVFRDLANHNQIFVKDKDPKVYNPNYYIDQVQLKDAQQT  
DLTSIQASFTTLDGVDKTEILKELKVTDKNQNAIQISDITLDTSKSLLIKG  
DFNPKQGHFNISYNGNNTTRQSWFKDQLYAYSGNLGAVLNQDGSKV  
EASLWSPSADSVTMIYDKDNQNRVVATTPLVKNNKGVWQITILDTKLG  
KNYTGYYLYEIKRGKDKVKILDPYAKSLAEWDSNTVNDDIKTAKAAF  
VNPSQLGPKNLSFAKIANFKGKQDAVIYEAHVRFDTSDQSLDGLKLNQL  
GTFAAFSEKLDYLQKLGVTHIQLLPVLSYFYVNEMDKSRSTAYTSSDNN  
YNWGYDPQSYFALSGMYSEKPKDPSARIAELKQLIHDIHGRGMGVIDV  
VYNHTAKTYLFEDIEPNYYHFMNEDGSPRESFGGGRLGTTHAMSRRLV  
VDSIKYLTSEFKVDGFRFDMMGDHDAAIELAYKEAKAINPNMIMIGEG  
WRTFQGDQGKPKPADQDWMKSTDTVGVFSDDIRNSLKSFPNEGTPA  
FITGGPQSLQGIFKNIKAQPGNFADSPGDVVQYIAAHDNLTLHDVIAKSI  
NKDPKVAEEDIHRRLRLGNMILTSQGTAFIHSGQEYGRTRKLLNPDYM  
TKVSDDKLPNKATLIEAVKEYPYFIHDSYDSSDAINHFDWAAATDNNKH  
PISTKTQAYTAGLITLRRSTDAFRKLSKAIDREVSLITEVGQGDIKEKDL

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FIG. 1 CONT'D

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VIAYQTIDSKGDIYAVFVNADSKARNVLLGEKYKHELLKGQVIVDADQA  
GIKPISTPRGVHFEKDSLIDPLTAIVIKVGKVAPSPKEELQADYPKTQSFK  
GSKTVEKVNRIANKTSITPVVSNKTD SYLTNEANLPKTGDKSSKILSVVG  
ISILASLLALLGLSLKRNRT\*

ID-6

Clone 9

ATGAAAAAAGTTTTTTTTCTCATGGCTATGGTTGTGAGTTTAGTAATGATAGCAGG  
GTGTGATAAGTCAGCAAACCCCAAACAGCCTACGCAAGGCATGTCAGTTGTAACC  
AGCTTTTACCCAATGTATGCGATGACAAAAGAAGTATCTGGAGACCTAAATGATGT  
GAGGATGATCCAATCAGGTGCAGGCATTCAATCCTTTGAACCGTCTGTAAATGATG  
TGGCAGCTATTTATGACGCGGATTTGTTTGTTTACCAATCACATACCTTAGAAGCTT  
GGGCAAGGGATCTAGACCCTAATTTAAAAAAATCAAAGGTTAATGTGTTTGAAGC  
GTCAAACCTCTGACACTAGATAGAGTCAAAGGGCTAGAAGATATGGAAGTCACA  
CAAGGCATTGACCCTGCGACACTTTATGACCCACATACCTGGACGGATCCCGTTTT  
AGCTGGTGAGGAAGCTGTTAATATCGCTAAAGAGCTAGGACATTTGGATCCTAAAC  
ACAAAGACAGTTACTATAAAAGGCTAAGGCTTTCAAAAAAGAAGCAGAGCAACT  
AACTGAAGAATACACTCAAAAATTTAAAAAGGTGCGCTCAAAAACATTTGTGACG  
CAACACACGGCATTCTTATCTGGCTAAACGATTCCGGCTTGAAACAACCTGGTAT  
CTCGGGTATTTCTCCAGAGCAAGAGCCCTCTCCTCGCCAATTGAAAGAAATCAAG  
ACTTTGTTAAAGAATACAACGTCAAGACTATTTTTGCAGAAGACAACGTCAACCCC  
AAAATTGCTCATGCTATTGCGAAATCAACAGGAGCTAAAGTAAAGACATTAAGTC  
CACTTGAAGCTGCTCCAAGCGGAAACAAGACATATCTAGAAAATCTTAGAGCAA  
TTTGAAAGTGCTCTATCAACAGTTGAAGTAA

MKKVFFLMAMVVSLVMIAGCDKSNPKQPTQGMSVVTSFYPMYAMTKEVSGDLND  
VRMIQSGAGIHSFEPVNDVAAIYDADLFVYQSHTLEAWARLDLDPNLKSKVNVFEAS  
KPLTLDRVKGLEDMEVTQGIDPATLYDPHTWTDVPLAGEEAVNIAKELGHLDPKHKD  
SYTKKAKAFKKEAEQLTEEYTKFKKVRSKTFVTQHTAFSYLAKRFLKQLGISGISPE  
QEPSRQLKEIQDFVKEYNVKTIFAEDNVNPKIAHAIAKSTGAKVKTLSPLEAAPSGNK  
TYLENLRANLEVLYQLK\*

ID-7

Clone 15

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TTGTTCAATAAAAATAGGTTTTAGAACTTGGAATCAGGAAAGCTTTG  
GCTTTATATGGGAGTGCTAGGATCAACTATTATTTTAGGATCAAGTCC  
TGTATCTGCTATGGATAGTGTTGGAATCAAAGTCAAGGTAATGTTTT  
AGAGCGTCGCCAACGTGATGCGGAAAACAAAAGTCAGGGTAATGTT  
TTAGAGCGTCGCCAACGTGATGCGGAAAACAAGAGCCAAGGCAATG  
TTTTAGAGCGTCGTC AACGCGATGTTGAGAATAAGAGCCAAGGCAAT

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FIG. 1 CONT'D

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GTTTTAGAGCGTCGTC AACGTGATGCGGAAAACAAAAGTCAGGGCA  
ATGTTCTAGAGCGCCGCAACGTGATGCGGATAACAAGAGCCAAGTA  
GGTCAACTTATAGGGAAAAATCCACTTTTTTCAAAGCCAAGTGTATCT  
AGAGAAAATAATCACTCTAGTCAAGGTGACTCTAACAAACAGTCATT  
CTCTAAAAAAGTATCTCAGGTTACTAATGTAGCTAATAGACCGATGT  
TAACTAATAATTCTAGAACAATTT CAGTGATAAATAAATTACCTAAA  
ACAGGTGGTGATCAAAATGTCATTTTTAACTTGTAGGTTTTGGTTTA  
ATTTTGTTAACAAAGTCGCTGCGGTTTGAGACGCAATGAAAATTA

MFNKIGFRTWKSGLWLYMGVLGSTIILGSSPVSAMDSVGNQSQGNVL  
ERRQRDAENKSQGNVLERRQRDAENKSQGNVLERRQRDVENKSQGNV  
LERRQRDAENKSQGNVLERRQRDADNKSQVGLIGKNPLFSKPTVSREN  
NHSSQGDSNKQSFSSKVSQVTNVANRPLMTNNSRTISVINKLPKTGGDQ  
NVIFKLVGFGLILLTSRCGLRRNEN\*

ID-8

Clone 17

ATGACAAAAAACTTATTATTGCTATATTAGCACTATGCACTATCTTAACCACTTCT  
CAAGCTGTTTTAGCTAAAGAAAAATCACAACTGTTACCATAAAAAACAACCTATTC  
GGTCTATATTAAGAAAAAGAAAAAGAGACAAGCCGGATAATAAAAAGCAAATCAG  
CGAGACACTTAAAGTTCCTTTAAAACCCAAAAAAGTAGTTGTTTTGATATGGGAG  
CTTTGGATACTATCACAGCTTTAGGAGCTGAAAAATCTGTTATTGGTATCCCGAAG  
GCTAAAAATGCTCTAAGTTTATTGCCCAATAACGTCAAATCTGTTTATAAAGCTAA  
GAGATACCAAGACGTAGGAAGTCTCTTCGAACCAAACCTTTGAAGCTATTGCTCGTA  
TGCAACCTGATGTGGTTTTCTAGGAGCACGTATGGCTTCTGTTGATAATATTGAA  
AAATTAAGGAGGCTGCACCTAAAGCAGCATTAGTATATGCTGGAGTCGACTCAA  
AAAAAGTATTTGACAAAGGAGTTGCTGAGCGTGTCACAATGTTAGGGAAAAATCTTC  
GACCAAAATAAAAAGGCAAAAACCTTTAATAAAGATATCGCACAAAGCTGTTCTTA  
AATTGCAGAAAACCTATTGAGAAAAAAGGTAAACCTACAGCTCTATTTGTAATGGC  
AAACAGCGGTGAACTTTTAACTCAATCACCTTCTGGTCGTTTTGGTTGGATTTTCTC  
TGTAGGTGGATTTAAAGCAGTCAATGAAAATGAAAACTAAGTTCACATGGTACTC  
CCGTATCTTATGAATACATCGCTGAAAAAAATCCTAACTATCTCTTTGTTTTAGATC  
GTGGAGCGACTATTGGACAAGGAGCTTCATCAAAGAAGCTTTTTAATAACGATGTT  
ATTAAGCAACTGATGCTGTCAAAAACAAACGTGTTTCATGAGGTAGATGGAAAAG  
ATTGGTATATCAATTCAGGCGGAAGCCGAGTAACACTCCGTATGATTAAAGATGTA  
CAGAACTTTGTTGATAATCGTTAA

MTKKLIILALCTILTTSQAVLAKEKSQTVTIKNNYSVYIKKEKRDKPDN  
KKQISETLVPLPKPKVVFDMGALDITITLGAEKSVIGIPKAKNALSLL  
PNNVKSVMYKAKRYQDVGSLFEPNFEIARMQPDVVFLGARMASVDNIE  
KLKEAAPKAALVYAGVDSKKVFDKGVAERVTMLGKIFDQNKKAKTFN  
KDIAQAVLKLQKTIEKKGKPTALFVMANSGELLTQSPSGRFGWIFSVGG

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FIG. 1 CONT'D

SUBSTITUTE SHEET (RULE 26)

FKAVNENEKLSSHGTPVSYEYIAEKNPNYLFLVLDRGATIGQGASSKELFN  
NDVIKATDAVKNRVHEVDGKDWYINSGGSRVTLRMIKDVQNFVDNR

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ID-9

Clone 18

GTGAAGAAAACATATGGTTATATCGGCTCAGTTGCTGCTATTTTACTAGCTACTCAT  
ATTGGAAGTTACCAGCTTGGTAAGCATCATATGGGTCTAGCAACAAAGGACAATC  
AGATTGCCTATATTGATGATAGCAAAGGTAAGGTAAGGTAAGGCCCTAAAACAAACAA  
AACGATGGATCAAATCAGTGCTGAAGAAGGCATCTCTGCTGAACAGATCGTAGTC  
AAAATTACTGACCAAGGTTATGTTACCTCACACGGTGACCATTATCATTTTTACAAT  
GGGAAAGTTCCTTATGATGCGATTATTAGTGAAGAGTTGTTGATGACGGATCCTAA  
TTACCATTTTAAACAATCAGACGTTATCAATGAAATCTTAGACGGTTACGTTATTA  
AAGTCAATGGCAACTATTATGTTTACCTCAAGCCAGGTAGTAAGCGCAAAAACATT  
CGAACCAAACAACAAATTGCTGAGCAAGTAGCCAAAGGA ACTAAAGAAGCTAAA  
GAAAAAGGTTTAGCTCAAGTGGCCCATCTCAGTAAAGAAGAAGTTGCGGCAGTCA  
ATGAAGCAAAAAGACAAGGACGCTATACTACAGACGATGGCTATATTTTTAGTCC  
GACAGATATCATTGATGATTTAGGAGATGCTTATTTAGTACCTCATGGTAATCACT  
ATCATTATATTCCTAAAAAAGATTTGTCTCCAAGTGAGCTAGCTGCTGCACAAGCC  
TACTGGAGTCAAAAACAAGGTCGAGGTTGCTAGACCGTCTGATTACCGCCCGACAC  
CAGCCCCAGGTCGTAGGAAAGCCCCAATTCTGATGTGACGCCTAACCTGGACA  
AGGTCATCAGCCAGATAACGGTGGTTATCATCCAGCGCCTCCTAGGCCAAATGATG  
CGTCACAAAACAAACACCAAAGAGATGAGTTTAAAGGAAAAACCTTTAAGGAACT  
TTAGATCATCTACACCGTCTTGATTTGAAATACCGTCATGTGGAAGAAGATGGGT  
TGATTTTTGAACCGACTCAAGTGATCAAATCAAACGCTTTTGGGTATGTGGTGCCT  
CATGGAGATCATTATCATATTATCCCAAGAAGTCAGTTATCACCTCTTGAAATGGA  
ATTAGCAGATCGATACTTAGCCGGCCAAACTGATGACAACGACTCAGGTTCAGATC  
ACTCAAACCATCAGATAAAGAAGTGACACATACCTTTCTTGGTCATCGCATCAAA  
GCTTACGGAAAAGGCTTAGATGGTAACCATATGATACGAGTGATGCTTATGTTTT  
TAGTAAAGAATCCATTCATTCAAGTGATAAATCAGGAGTTACAGCTAAACACGGA  
GATCATTTCCTACTATATAGGATTTGGAGA ACTTGAACAATATGAGTTGGATGAGGT  
CGCTAACTGGGTGAAAGCAAAGGTCAAGCTGATGAGCTTGTGCTGCTTTGGATC  
AGGAACAAGGCAAAGAAAAACCACTCTTTGACACTAAAAAAGTGAGTCGCAAAGT  
AACAAAAGATGGTAAAGTGGGCTATATTATGCCAAAAGATGGCAAGGACTATTTCT  
TATGCTCGTTATCAACTTGATTTGACTCAGATTGCCTTTGCCGAACAAGAACTAATG  
CTTAAAGATAAGAAGCATTACCGTTATGACATTGTTGATACAGGCATTGAGCCACG  
ACTTGCTGTAGATGTGTCAAGTCTGCCGATGCATGCTGGTAATGCTACTTACGATA  
CTGGAAGTTCGTTTGTTATCCACATATTGATCATATCCATGTCGTTCCGTATTCAT  
GGTTGACGCGCAATCAGATTGCAACAATCAAGTATGTGATGCAACACCCCGAAGT  
TCGTCCGGATGTATGGTCTAAGCCAGGGCATGAAGAGTCAGGTTCCGGTCATTCCAA  
ATGTTACGCCTCTTGATAAACGTGCT

FIG. 1 CONT'D



GGTATGCCAAACTGGCAAATTATCCATTCTGCTGAAGAAGTTCAAAAAGCCCTAGC  
 AGAAGGTCGTTTTGCAGCACCCAGACGGCTATATTTTCGATCCACGAGATGTTTTGG  
 CAAAAGAACTTTTTGTATGGAAAGATGGCTCCTTTAGCATCCCAAGAGCAGATGGC  
 AGTTCATTGAGAACCATTAATAAATCCGATCTATCCCAAGCTGAGTGGCAACAAGC  
 TCAAGAGTTATTGGCAAAGAAAAATGCTGGTGATGCTACTGATACGGATAAACCT  
 GAAGAAAAGCAACAGGCAGATAAGAGCAATGAAAACCAACAGCCAAGTGAAGCC  
 AGTAAAGAAGAAAAAGAATCAGATGACTTTATAGACAGTTTACCAGACTATGGTC  
 TAGATAGAGCAACCCTAGAAGATCATATCAATCAATTAGCACAAAAAGCTAATAT  
 CGATCCTAAGTATCTCATTTTCCAACCAGAAGGTGTCCAATTTTATAATAAAAATG  
 GTGAATTGGTAACTTATGATATCAAGACACTTCAACAAATAAACCTTAA

MKKTYGYIGSVAAILLATHIGSYQLGKHHMGLATKDNQIAYIDDSKGGKVKAPKTNKT  
 MDQISAEEGISAEQIVVKITDQGYVTSBGDHYHFYNGKVPYDAIISEELMTDPNYHFK  
 QSDVINEILDGYVIKVNNGNYVYLKPGSKRKNIRTKQQIAEQVAKGTKEAKEKGLAQV  
 AHLKEEVAAVNEAKRQGRYTTDDGYIFSPTDIIDDLGDAYLVPHGNHYHYIPKKDLS  
 PSELAQAAYWSQKQGRGARPSDYRPTAPGRRKAPIPDVTPNPGQGHQPDNGGYHP  
 APPRPNDASQNKHQRDEFKGTKFKELLDHLHRLDLKYRHVEEDGLIFEPTQVIKSNAF  
 GYVVPBGDHYHIIPRSQLSPLEMELADRYLAGQTDNDSDGSDHSPSKDKEVTHFTLGH  
 RIKAYGKGLDGKPYDTSDAYVFSKESIHSVDKSGVTAKHGDHFHYIGFGELEQYELDE  
 VANWVKAKGQADELVAALDQEQGKEKPLFDTKKVSARKVTKDGKVGYPKDGKDY  
 FYARYQLDLTQIAFAEQELMLKDKKHRYRDIIVDTGIEPRLAVDVSSSLPMHAGNATYD  
 TGSSFVIPHIDHIHVVPYSWLTRNQIATIKYVMQHPEVRPDVWSKPGHEESGSVIPNVTP  
 LDKRAGMPNWQIIHSAEEVQKALAEGRFAAPDGYIFDPRDVLAKETFVWKDGSFSIPR  
 ADGSSLRTINKSDLSQAQEWQQAQELLAKKNAGDATDTDKPEEKQQADKSNENQQPSE  
 ASKEEKESDDFIDSLPDYGLDRATLEDHINQLAQKANIDPKYLIFQPEGVQFYNGKNGEL  
 VTYDIKTLQQINP\*

ID-10

Clone 22

ATGATACGCCAGTTTTTAAGAGAACACTTGATTTGGTATATTTTATATATCATGATG  
 TTTGTCCTATTTTTTATTAGTTTCTATCTATATCATTTACCAATGCCCTATTTGTTA  
 ATTCCTTAGGTTTAAATGTTATTGTTTTACTAGGAATTAGTATTTGGCAATACAGTC  
 GTTACAGGAAAAAATGTTACATCTCAAATATTTAATAGTAGTCAGGACCCCTCT  
 TTCGAACTTCAACCGAGTGATTACGCTTATTTAATATTATTACACAATTAGAAGCT  
 AGAGAAGCGCAAAAAGTTTCTGAAACAATTGAACAAACCAATCATGTTGCACTTA  
 TGATAAAGATGTGGTCGCACCAAATGAAAGTTCCATTGGCAGCTATTTCAATTAATG  
 GCCCAGACAAATCATCTCGATCCTAAGGAAGTTGAACAACAATTATTGAAATTGCA  
 ACATTATCTTGAAACGTTGTTAGCATTTTTGAATTTAGACAATATCGTGACGATTT  
 TCGTTTTGAAGCTGTTAGCCTTAGAGAAGTAGTAGTAGAAATTATAAAATCGTATA  
 AGGTTATTTGTCTATCCAAAAGCTTATCTATCATAATTGAAGGCGATAATATCTGG  
 AAAACAGACAAAAAGTGGTAACTTTTGCTCTTTCACAGGTGCTAGATAATGCCAT

FIG. 1 CONT'D

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AAAATATTCTAATCCTGAGTCAAAGATAATAATAAGCATAGGAGAAGAGAGTATT  
 AGAATACAAGACTACGGTATCGGCATACTCGAAGAGGATATCCCTAGACTTTTTGA  
 AGATGGCTTTACGGGTTACAACGGTCATGAGCACCAAAGGCAACAGGCATGGGG  
 TTATATATGACAAAAGAAGTCTTATCTAGTCTGAATTTGTCCATTTCCGGTGGATAGC  
 AAAATTAATTATGGGACTGCTGTTTCTATACATAAATAA

MIRQFLREHLIWYILYIMMFVLFISFYLYHLPMPYLFNSLGLNVIVLLGISIWQYSRYR  
 KKMLHLKYFNSSQDPSFELQPSDYAYFNITQLEAREAQKVSETIEQTNHVALMIKMW  
 SHQMKVPLAAISLMAQTNHLDPKEVEQQLKLOHYLETLLAFLKFRQYRDDFRFEAV  
 SLREVVVEIKSYKVICLSKSLSIIEGDNIWKTDKKWLTFALSQVLDNAIKYSNPESKIIIS  
 IGEESIRIQDYGIGILEEDIPRLFEDGFTGYNGHEHQKATGMGLYMTKEVLSSLNLSISV  
 DSKINYGTAVSIHKZ

ID-11

Clone 23

ATGACTTATCAAAAAACAGTTGTTTTGGCTGGTGATTATTCCTACATTAGACAAATT  
 GAAACCACATTAATAATCTCTCTGTGTCTATCATGAGAATCTCTCAATTTTTATTTTT  
 AATCAAGATATTCCTCAAGAATGGTTTTTAGCTATGAAAGATAGGGTTGGACAAAC  
 TGGAATCAAATTCAGGATGTAAAGCTCTTCCATGATCACTTATCCCCAAAATGGG  
 AAAATAAAAAGCTTAATCATATTAATTATATGACCTATGCTCGTTATTTACATACCTC  
 AGTACATCTCAGCTGATACAGTTTTATATCTTGACTCTGACTTAGTTGTTACTACTA  
 ATTTAGATAACCTCTTTCAAATTTCACTAGACAATGCATATTTAGCTGCAGTTCAG  
 CTCTTTTTGGGCTTGGATATGGGTTTAATGCTGGAGTAATGGTAATTAACAACCAA  
 CGTTGGCGACAAGAAAATATGACTATTAATTAATTGAAAAAATCAAAAGGAAA  
 TTGAGAATGCCAACGAAGGGGATCAAACAATTCTTAATCGCATGTTTGAAAATCAG  
 GTAATTTATTTAGATGATACCTACAATTTTCAAATTTGGTTTTGATATGGGAGCTGCT  
 ATCGATGGGCATAAATTTATTTTTGACATCCCAATTACCCCACTCCCAAAAATTATT  
 CACTACATTTCCGGGAATCAAACCTTGGCAAACATTATCAAATATGAGACTCCGTGA  
 GGTATGGTGGCACTATAATTTACTTGAATGGTCAAGTATCATATCTAGTAAAAAAG  
 TATTTGGTTTAGACCACCCAATTAACACAAAATTATCGTCTCAATTTCCTTATTG  
 CTACAACCTTCTGATTGTATACCATCTATCTCAGAATTAGTCACTGCCCTTCCAGATT  
 GTCTATTTACATTGCATGCACCAACAGTTATGTCTGA

MTYQKTVVLAGDYSYIRQIETTLKSLCVYHENLSIFIFNQDIPQEWFLAMKDRVGQTG  
 NQIQDVKLFHDHLSPKWENKLNHNINMTYARYFIPQYISADTVLYLDSDLVVTNLD  
 NLFQISLDNAYLAAVPALFGLGYGFNAGVMVINNRWRQENMTIKLIEKNQKEIENAN  
 EGDQITLNRMFENQVIYLD DTYNFQIGFDMGAAIDGHKFIFDIPITPLPKIIHYISGIKPW  
 QTLNMLREVWWHYNLLEWSSISSKKVFLGDHPIKTQNYRLNFLIATTSDCIPSISEL  
 VTALPDCLFHIACTNSYV\*

ID-12

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FIG. 1 CONT'D

Clone 27

GTGAAGAAAACATATTGTTATATCGGCTCAGTTGCTGCTATTTTACTAGCTACTCAT  
 ATTGGAAGTTACCAGCTTGGTAAGCATCATATGGGTCTAGCAACAAAGGACAATC  
 AGATTGCCTATATTGATGATAGCAAAGGTAAGGTAAAAGCCCCTAAAACAAACAA  
 AACGATGGATCAAATCAGTGCTGAAGAAGGCATCTCTGCTGAACAGATCGTAGTC  
 AAAATTACTGACCAAGGTTATGTTACCTCACACGGTGACCATTATCATTTTTACAAT  
 GGGAAAGTTCCTTATGATGCGATTATTAGTGAAGAGTTGTTGATGACGGATCCTAA  
 TTACCATTTTAAACAATCAGACGTTATCAATGAAATCTTAGACGGTTACGTTATTA  
 AAGTCAATGGCAACTATTATGTTTACCTCAAGCCAGGTAGTAAGCGCAAAAACATT  
 CGAACCAACAACAATTGCTGAGCAAGTAGCCAAAGGAACTAAAGAAGCTAAA  
 GAAAAAGGTTTAGCTCAAGTGGCCCATCTCAGTAAAGAAGAAGTTGCGGCAGTCA  
 ATGAAGCAAAAAGACAAGGACGCTATACTACAGACGATGGCTATATTTTLAGTCC  
 GACAGATATCATTGATGATTAGGAGATGCTTATTTAGTACCTCATGGTAATCACT  
 ATCATTATATTCCTAAAAAAGATTTGTCTCCAAGTGAGCTAGCTGCTGCACAAGCC  
 TACTGGAGTCAAAAACAAGGTCGAGGTGCTAGACCGTCTGATTACCGCCCGACAC  
 CAGCCCAGGTCGTAGGAAAGCCCACTTCTGATGTGACGCCTAACCTGGACAA  
 GGTCATCAGCCAGATAACGGTGGTTATCATCCAGCGCCTCCTAGGCCAAATGATGC  
 GTCACAAAACAACACCAAAGAGATGAGTTTAAAGGAAAAACCTTTAAGGAACTT  
 TTAGATCAACTACACCGTCTTGATTTGAAATACCGTCATGTGGAAGAAGATGGGTT  
 GATTTTGAACCGACTCAAGTGATCAAATCAAACGCTTTTGGGTATGTGGTGCCTC  
 ATGGAGATCATTATCATATTATCCCAAGAAGTCAGTTATCACCTCTTGAAATGGAA  
 TTAGCAGATCGATACTTAACCCGGCCAAACTGA

MKPTYCYIGSVAAILLATHIGSYQLGKHHMGLATKDNQIAYIDDSKKGKVKAPKTNKT  
 MDQISAEEGISAEQIVVKITDQGYVTSHGDHYHFYNGKVPYDAIISEELMTPNYHFK  
 QSDVINEILDGYVIKVNNGNYVYLKPGSKRKNIRTKQQIAEQVAKGTKEAKEKGLAQV  
 AHLKKEEVAAVNEAKRQGRYTDDGYIFSPDIIDDLGDAYLVPHGNHYHYIPKDL  
 PSELAQAQYWSQKQGRGARPSDYRPTAPGRRKAPLPDVTNPNGQGHQPDNGGYHP  
 APPRPNDASQNKHQRFDEFKGTKFKELLDQLHRLDLKYRHVEEDGLIFEPTQVIKSNF  
 GYVVPBGDHYHIIPRSQLSPLEMELADRYLTRPN\*

ID-13

Clone 28

ATGGTAAATGATATATTAGAAAGAATGTATAAAGAGAATATTCCAAAATCTTACCT  
 TACATCCGTCCCATTAGTTATTTCTCAAAAAGGAAGAACAACCTATTCGTTTAGTAT  
 GACTGGTGGTCAACAAATAGATGGAGTGAAATTCACACAGATATATGAGGACTAT  
 ATGAAATTACTCAGTCAAGGTAAGGATATCGCAGAGTTATATCAAAAATATTCTAA  
 AGAAGAGTTGGCAAATCTAGGCATTAATATTTATCAATCCAATGATATAGAAAGG  
 ACTGAGGAAAGAACTTTTGATGAAATTATCAGTTGGGTTTCCAACCCTTATGCAAC  
 AAGACCAATTCAAGAAAGGCACACTATTCAATTAGAGCCAACAAGATTTTCACTA

FIG. 1 CONT'D

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GAGGATAAGAAAAGAATTGAAGAAGCTGCAGCTCAAGGACTAAGCGAAATCGAC  
 CTTATTGATTTAGTTGACCTATATGATATTAATTTAGACAATACAAGCGTCAATCGC  
 CATATTGTGGGGTTATTGACTAATAACACCCAAGTAACATACTATTTCCAAGAACA  
 ATTAAATAAGGAGTTGCTGTCAATGGCTCACGCTTTAGATAACGTACAACAGGCCT  
 TTATTAAATTATTAAGTGAAGAGGAGATACGAAAATTTGCTCTTTAA

MVNDILERMYKENIPKSYLTSVPLVISQKGRTTYFSMTGGQQIDGVKFTQIYEDYMK  
 LLSQGKDIAELYQKYSKEELANLGINIYQSDIERTEERTFDEIISWVSNPYATRPIQERH  
 TIQLEPTRFSLEDKKRIEAAAQGLSEIDLIDLVDLYDINLDNTSVNRHIVGLLTNNTQV  
 TYYFQEQLNKELLSMAHALDNVQQAFIKLLSEEEIRKFAL\*

ID-14

Clone 31

ATGAATAAAAGAAGAAAATTATCAAAATTGAATGTAAAAAACAACATTTAGCTT  
 ATGGAGCTATCACTTTAGTAGCCCTTTTTTTCATGTATTTTGGCTGTAACGGTCATCT  
 TAAAAGTTCACAAGTTACTACTGAATCTTTGTCAAAAGCAGATAAAGTTCGCGTA  
 GCCAAAAAATCAAAAATGACTAAGGCGACATCTAAATCAAAAGTAGAAGATGTAA  
 AACAGGCTCCAAAACCTTCTCAGGCATCTAATGAAGCCCCAAAATCAAGTTCTCAA  
 TCTACAGAAGCTAATTCTCAGCAACAAGTTACTGCGAGTGAAGAGGCGGCTGTAG  
 AACAAGCAGTTGTAACAGAAAATACCCCTGCTACCAGTCAGGCACAACAACTTA  
 TGCTGTTACTGAGACAACTTACAAACCTGCTCAACACCAGACAAGTGGCCAAGTAT  
 TGAGCAATGGAAATACTGCAGGGGCGGTTCGGATCTGCTGCTGCAGCACAAATGGC  
 TGCTGCAACAGGAGTCCCTCAGTCTACTTGGGAACATATTATTGCCCGTGAATCAA  
 ATGGTAATCCTAATGTTGCTAATGCCTCAGGGAGCTTCAGGACTTTTCCAAACGAT  
 GCCAGGTTGGGGTTCAACAGCTACAGTTCAGGATCAAGTTAA

MNKRRLSKLNVKKQHLAYGAILLVALFSCILAVTVIFKSSQVTTESLSKADKVRVAK  
 KSKMTKATSKSKVEDVKQAPKPSQASNEAPKSSSQSTEANSQQQVTASEEAAVEQAV  
 VTENTPATSQAQQTAVTETTYKPAQHQTSGQVLSNGNTAGAVGSAAAQMAAATG  
 VPQSTWEHILARESNGNPVANASGASGLFQTMPGWGSTATVQDQVNSAIKAYRAQG  
 LSAWGY\*

ID-15

Clone 32

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ATGATTGTTGGACACGGAATTGATTTACAAGAGATAGAGGCGATTACTAAAGCAT  
 ATGAGCGTAATCAACGTTTTTGCAGAACGCGTTTTGACCGAACAAGAATTGCTTCTT  
 TTAAAGGAATTTCCAATCCCAAGCGTCAGATGTCTTTTTTAACAGGGCGATGGGC  
 AGCAAAAGAGGCTTATAGCAAAGCACTTGGAACAGGAATTGGGAAAGTTAATTTT  
 CATGATATCGAAATTTTATCGGATGATAAAGGAGCGCCTTTGATTACAAAAGAACC

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FIG. 1 CONT'D

---

GTTTAATGGAAAATCTTTTGTTCATATCTCATAGTGGTAATTATGCACAAGCTAG  
TGTTATTTTGGAGGAAGAAAAATGA

MIVGHGIDLQEIEAITKAYERNQRFAERVLTEQELLFLKGISNPKRQMSFLTGRWAAKE  
AYSKALGTGIGKVNFDIEILSDDKGAPLITKEPFNGKSFVSISHSGNYAQASVILEEEK\*

ID-16

Clone 35

ATGATTTTTGTACAGTGGGGACACATGAACAGCAGTTCAACCGTCTTATTAAGA  
AGTTGATAGATTAAGGGACAGGTGCTATTGATCAAGAAGTGTTTCATTCAAACG  
GGTACTCAGACTTCGAACCTCAGAATTGTCAGTGGTCAAATTTCTCTCATATGAT  
GATATGAACTCTTACATGAAAGAAGCTGAGATTGTTATCACACATGGCGGCCCAGC  
GACGTTTATGTCAGTTATTTCTTTAGGGAAATTACCAGTTGTTGTTCTTAGGAGAAA  
GCAGTTTGGTGAACATATCAATGATCATCAAATACAATTTTTAAAAAAATTGCC  
ACCTGTATCCCTTGGCTTGGATTGAAGATGTAGATGGACTTGCGBAAGCGTTGAAA  
AGGAATATAGCTACAGAAAAATATCAGGGAAATAATGATATGTTTTGTCATAAATT  
AGAAAAAATTATAGGTGAAATATGA

MIFVTVGTHEQQFNRLIKEVDRLKGTGAIDQEVFIQTGYSDFEPQNCQWSKFLSYDDM  
NSYMKEAEIVITHGGPATFMSVISLGLPVVPRRKQFGEHINDHQIQFLKKIAHLYPL  
AWIEDVDGLAEALKRNIAATEKYQGNNDMFCHKLEKIIGEI\*

ID-17

Clone39

TTGGAAGACAAATTATTCAACAAACATTTTATAGGCATTACTATTTTAACTTTATT  
GTTTATATGGTCTATTATTTGTTACCGTTATCATAGCTTTTATTGCGACTAAAGAG  
TTAGGTGTTAGCACTAGCCAAGCAGGATTAGCAACGGGGATTATATTGTAGGGAC  
TTTGATTGCTCGTCTTATATTTGGTAAGCAATTAGAAGTTCTAGGACGTAAGTTAGT  
TTTACGTGGAGGGGCTATTTTTACTTACTAACAACCTTAGCTTATTTTTATATGCC  
AAGTATCGGAGTAATGTATTTAGTTCGTTTCCTAAATGGTTTTGGTTATGGCGTCGT  
GTCAACAGCAACTAATACTATTGTAACAGCCTATATAACCAGCTGATAAAAGAGGTG  
AGGGGATTAACCTTTACGGTCTATCAACAAGTTTAGCCGCAGCTATTGGTCCTTTG  
TAGGAACATTTATGCTAGACAACCTTCATATTAACCTTAAAATGGTTATTGTATTAT  
GTAGTATTTAATTGCGATTGTAGTGTGGGAGCATTGTTTTCCAGTCAAAAATA  
TTACTTTAAATCCAGAACAGTTAGCTAAATCAAATCATGGACTATTGATAGTTTC  
ATTGAGAAAAAAGCAATTTTTATCACAATTATTGCATTTTGTATGGGTATCTCCTAT  
GCTCCGTGTTAGGTTTCCAAAAATTATATAACAAGAAATTAATTTGATGACAGT  
AGGAGCTTATTTCTTTATTGTTTATGCACTTGTCATCACTTAACCAGACCATCTAT  
GGGAAGATTAATGGACGCTAAGGGAGATAAGTGGGTGCTTTATCCAAGTTATCTGT  
TCTTAACCTTTGGGACTTGCTTTATTAGGGAGTGCTATGGGAAGTGTTACCTACCTTC

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FIG. 1 CONT'D

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TATCAGGTGCTTTGATTGGTTTTGGTTATGGCACCTTTATGTCTTGTGGCCAAGCAG  
 CATCAATCAAAGGTGTTGAGGAACATCGTTTCAATACAGCCATGTCAACTTACATG  
 ATAGGTCTTGATTTAGGGTTAGGTGCTGGACCTTACATTTTGGGACTTGTTAAAGAT  
 GGTTTTCTTGGAGCTGGTGTGCAATCCTTTAGAGAATTATTCTGGATAGCAGCGATT  
 ATCCTGTTGTTTGTGGTATTCTATATTTCTTAAAATCATCTAGACAAGTTGAAACT  
 AAAACTATA  
 TAA

MEDKLFNKHFIGITILNFIVYMVYYLFTVIIAFIATKELGVSTSQAGLATGIYIVGTLIARL  
 IFGKQLEVLGRKLVLRGGAIFYLLTTLAYFYMP SIGVMYLVRFLNGFGYGVVSTATNTI  
 VTAYIPADKRGEINFYGLSTSLAAAIGPFVGT FMLDNLHINFKMVIVLCSILIAIVVLG  
 AFVFPVKNITLNPEQLAKSKSWTIDSFIEKKAIFITIIAFLMGISYASVLGFQKLYTTEINL  
 MTVGAYFFIVYALVITLTRPSMGR LMDAKGDKWVLYPSYLFLTLGLALLGSAMGSVT  
 YLLSGALIGFGYGT FMSCGQAASIKGVEEHRFNTAMSTYMIGLDLGLGAGPYILGLVK  
 DGFLGAGVQSFRELFWIAAIPVVC GILYFLKSSRQVETKTIZ

ID-18

Clone 47

ATGAATAGTGAACCTAAAAGTCAGTCAAACGAAGTAAAAAATAGCAAGCAATCAG  
 AAGTGAAGAAAGATAAAAAAATGACAAAAAAGAACAATTAGCCTATCTCAAAG  
 AGCATGAGCAAGAAATCATAGATTATGTAAAATTACATAACAACCAAATTGAGTC  
 CGTTCAATTTCGATTGGTCAAGTGTAAGTAGAACAAGCGGGAATGGAACTCCA  
 CAAGGGGGTGATTATAATCTTTCACTGAGAGGAAAGTTAATCATCTACAAAATTC  
 AAAATTAATAGTTGATTTTTATTTAGCTCATAAAAATGATATCCCAAATATCAAAT  
 CAATGGGAATGCTAATAAGCCATATATACATAAAAATGGTATTTGGCACATTTAT  
 GAATAG

MILGGCQMNSEPKSQSNEVKNSKQSEVKKDKKMTKKEQLAYLKEHEQEIIDYVKLHN  
 NQIESVQFDWSSVKVEQSGNGTPQGGDYNLSLRGKFNHLQNSKLIVDFYLAHKNDIPN  
 IKSMGMLNKPYIHKNGIWHIYEZ

ID-19

Clone 102

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ATGAAAAGATTTCGATTATCAAAGTTTATTA AAAATGATTGTTGTTATTTTGTTTT  
 ATTAGTGTAGCAGCTAGTTTTTATTTTTTCCACGTTGCCCAAGTTCGAGATGATAAA  
 TCCTTTATTTCAAATGGTCAACGTAAGCCTGGAAACTCTTATATGCTTATGATAAA  
 TCCTTTGATAAGCTATTAAGCAAAAAATAGAAATGACAAACCAAATATAAAGC  
 AAGTTGCTTGGTATGTTCTGCTGCTAAGAAA ACTCATAAGACAGTTGTTGTCGTTT  
 ATGGTTTTGCGAATAGCAAAGAGAATATGAAGGCATATGGTTGGCTGTTTCATAAG  
 TTAGGATACAATGTTCTTATGCCTGACAACATTGCACATGGTGAAAGTCATGGGCA

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FIG. 1 CONT'D

GTTGATAGGCTATGGCTGGAACGACCGCGAGAACATTATCAAATGGACAGAAATG  
 ATAGTGGATAAGAATCCATCAAGCCAAATTACTTTATTTGGTGTTCATGGGTGG  
 AGCAACAGTCATGATGGCTAGTGGTGAAAAATTACCTAGTCAGGTTGTTAATAT  
 CATTGAAGATTGTGGTTATTCTAGTGTGGGATGAATTAATAATTCAGGCTAAAG  
 AGATGTATGGTTTACCAGCCTTCCCCTTATATGAAGTTTCAACAATTTCTAAAA  
 TCAGAGCAGGTTTTTCGTATGGACAAGCAAGTAGTGTCGAACAATTGAAAAAGAA  
 TAATTTACCAGCCCTCTTTATTCATGGTGATAAGGATAATTTTGTTCACAACAAGTAT  
 GGTATATGACAATAAAGCTACAGCAGGTAAGAAAGAGCTTTATATTGTAAAA  
 GGGGCAAACATGCGAAATCTTTTGAACAGAGCCAGAAAAATATGAGAAACGTA  
 TCTCTAGTTTTTTGAAAAAATATGAAAAATAA

MKKIRLSKFIKMIVVILFLISVAASFYFFHVAQVRDDKSFISNGQRKPGNSLYAYDKSFD  
 KLLKQKIEMTNQNIKQVAWYVPAKKTHKTVVVVHGFANSKENMKA YGWL FHKLG  
 YNVLMPDNIAHGESHGQLIGY GWNDRNIKWTEMIVDKNPSSQITLFGVSMGGATV  
 MMASGEKLP SQV VNIIEDCGYSSVWDELKFQAKEMYGLPAFPLL YE VSTISKIRAGFSY  
 GQASSVEQLKKNL PALFIHGDKDNFVPTSMVYDNYKATAGKKEL YIVKGAKHAKSF  
 ETEPEKYEKRISSFLKKYEK\*

ID-20

Clone 120

TTGAGGAGTAATATGGTAAAGACAGCAGTTTTAATGGCGACATACAATGGCGAAA  
 AATTTATATCTGAACAACCTTGATTCAATTCGCCAACAGACATTA AAA ACCAGATTAT  
 GTATTATTGAGGGGATGATTGTTCAACGGATGAAACAGTCAATGTCGTCAATAACTA  
 TATCGCAAACATGAGTTAGAAGGCTGGAAAATTGTTAAAAACGACAAAAACTTA  
 GGCTGGCGTTTTAAATTTTCGTCAATTACTTATTGATGTGTTAGCCTATGAGGTTGAC  
 TATGTCTTTTTTAGTGATCAAGATGATATTTGGTATCTTGATAAAAACGAACGACA  
 GTTTGCCATTATGTCAGATAACCCTCAAATTGAGGTTTTGAGTGCAGACGTTGATA  
 TCAAACGATGTCTACAGAAGCCAGTGTTCCACATTTTCTAACTTTTTCTTCTAGTG  
 ATAGAATCAGTCAGTATCCTAAAGTATATGATTATCAAACATTCCGTCCCGGATGG  
 ACCATTGCTATGAAGAGAGATTTTGC GCAAGCTATCGCTTGA

MRSNMVKTAVLMATYNGEKFISEQLDSIRQQTLKPDYVLLRDDCSTDETVNVVNNYI  
 AKHELEGWKIVKNDKNL GWRLNFRQLLIDVLA YEVDYVFFSDQDDI WYLDKNERQF  
 AIMSDNPQIEVLSADVDIKTMSTEASVPHFLT FSSSDRISQYPKVYDYQTFRPGWTIAM  
 KRDFQAIAZ

ID-21

Clone 143

ATGATTCATGAGATTCACGATTGTCAATTTATTGAAAAAGGAAGTTACGTTTATTT  
 GAATTATATTAATGCTGAGGGCGAGAGAGTAGTTATTATAATCATAGATTTTGTCC

FIG. 1 CONT'D

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GTAGTGTAGTCCTATTTTATATCGTCTATTTATGATTTTACTTGCACAAGAAGTAC  
CTCACTTGCATGATTACATCTATAATGCAAGAGATGATCACTACGATACTTGGAAG  
TTTAAAGAATTAAGGAGTCAAACCATCCAGTCCTTTTGGCATTCTCTGAAAGGTG  
GCACGATAGTCGCTTGACTTCTAAAAGCCTTGCAGAATGTTTACAATTAACCGACC  
TTGATGAAGAAGTGAAATCGACCATCATTCAATTAAGACAGTTCGAAAAATCAGTC  
AGAAATCCTTTGGCTCACCTGATTAACCTTTTATGATGAGCAAGAACTATATCGTAC  
AACTCAATTTTCTTCTCAAGCATTTTATAGACCAGATTATCTTCTTGGCAAAGGTAAT  
TGGTGTTGAGTATGATACTGTAAATTTTCACTACGATACGGTTAACAAGCTTATTAT  
AAAGATACTTGAGTAA

MIHEIHDCQFIEKGSYVYLNINAEGERVVIIIIDFVRSVSPILYRLFMILLAQEVPHLHD  
YIYNARDDHYDTWKFKELKESNHPVLLAFSERWHDSRLTSKSLAECLQLTDLDEEVKS  
TIIQLRQFEKSVRNPLAHLIKPFDEQELYRTTQFSSQAFLDQIIFLAKVIGVEYDTVNFHY  
DTVNKLIKILE\*

ID-22

Clone 1

ATGGTAAAAGTTTCAAATTTAGGGTATCCACGTCTTGGTGAACAGCGCGAATGGAA  
GCAAGCGATCGAAGCTTTCTGGGCAGGGAATCTTGAACAAAAAGATTTAGAAAAA  
CAACTAAAACAATTACGTATCAATCATTTAAAGAAACAAAAAGAGGCAGGTATTG  
ACCTTATTCCAGTGGGGGATTTTCTTGTATGATCATGTTTTGGATTGTTCATTTCA  
ATTCAATGTAATCCCAAAGCGTTTCGATGAGTATGAGAGGAATTTAGACCTTTATT  
TTGCTATTGCAAGAGGTGACAAAGATAATGTCGCATCATCTATGAAAAAGTGGTTT  
AATACCAACTACCACTACATAGTCCCAGAATGGGAGGTTGAGACTAAACCTCACTT  
GCAGAATAATTACTTACTTGATCTTTATCTAGAAGCTAGGGAAGTAGTTGGTGATA  
AAGCAAAGCCGGTTATC

MEEIMVKVSNLGYPRLGEQREWKQAIEAFWAGNLEQKDLEKQLKQLRINHLKKQKE  
AGIDLIPVGFDFSCYDHVLDLSFQFNVIPKRFDEYERNLDLYFAIARGDKDNVASSMKK  
WFNTNYHYIVPEWEVETKPHLQNNYLLDLYLEAREVVGDKAKPVI

ID-23

Clone 2

ATGGTGTTACTTTTATTGCTAATGGTAGCCAAGTCAAGTTTGTGTTACATGGCTG  
TTTATAACGATACTGACAAAAATAAAATGTTACCAGATATGGAGGAAGGAGAAAG  
TTATCAAGTTAA

MVLLLLLMVAKSSLMVTWLFITILTKIKCYQIWRKEKVIKL

ID-24

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FIG. 1 CONT'D



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 Clone 14

ATGAACAAAAAATTTCGGGATCGGCTTGGCTTCGATTGCAGTACT  
 TAGTTTAGCTGCATGTGGACATCGTGGTGCCTTCTAAATCTGGTGGTAA  
 ATCAGATAGCTTGAAGGTTGCAATGGTAACAGATACCGGTGGTGTG  
 ATGATAAATCATTAAACCAATCTGGTTGGGAAGGTATGCAAGCTTGG  
 GGCAAGAAGAATGGCCTTAAAAAAGGAGCTGGTTTTGACTATTTCCA  
 ATCGGCAAGTGAATCTGATTATGCAACTAACTTAGATACAGCTGTGT  
 CTAGTGGTTATAAATTGATTTTCGGTATTGGATTTTCTCTTCATGATG  
 CTATTGATAAAGCAGCAGACAATAACAAAGATGTTAATTACGTCATC  
 GTTGATGATGTTATTAAGGGAAAGATAATGTTGCAAGTGTTGTCTTT  
 GCGGATAATGAATCAGCTTACTTAGCAGGTATTGCAGCCGCTAAAAC  
 TACCAAAACAAAAACAGTTGGCTTTGTAGGTGGTATGGAATCTGAGG  
 TTATTACCCGTTTTGAAAAAGTTTTGAAGCAGGTGTCAAATCAGTTG  
 ATAAATCAATAAAATTAAGTTGACTATGCTGGTTCATTCCGGTGAT  
 GCTGCTAAGGGTAAGACAATTGCAGCCGCACAATATGCTTCTGGCGC  
 AGATATT

MNKKISGIGLASIAVLSLAACGHRGASKSGGKSDSLKVAMVTDGTVGVD  
 DKSFNQSGWEGMQAWGKKNGLKKGAGFDYFQSASESDYATNLDTAVS  
 SGYKLIFGIGFSLHDAIDKAADNNDKDVNYVIVDDVIKGDNDVASVVFAD  
 NESAYLAGIAAAKTTKTKTVGFVGGMESEVITRFEKGFVKSVDKSI  
 KIKVDYAGSFGDAAKGTIAAAQYASGADI

## ID-25

## Clone 20

ATGTTACATTCTAAAAAATACATTCCTTATCGCTTATTGCCGTTCTC  
 TCTTTAGCAACATATACGAGTTTACAACCAAATCATGTAGCGGCTGA  
 ACAATCACAAAAACATCAACTGTTCTTATGAGTCAAAAAACTATTG  
 AACATAAGTTAAAAGTTGCAGATAAAGAAGCTGCTCCTCTCTACGCT  
 AAAATCGACCATATCCAACGACATATTGAAGTCAAAAAAGCAAAG  
 ATTTAAAAGTTATTGAATTGTATATTAACAAAGATATCAACCAACTA  
 GAGAAGCAAATAAACGTCTACTAACTAAATTCTATACTTCTATTGA  
 TAATCAAACATGGGATAGCACAAGTGAAGTCAAAAAATTGATTGATA  
 AGACAACCCTATCCACTAACGAAAAAGATAGATTAATAATTATTTTT  
 GAACAACGTGCTTACCTTGAGACAAGGTTGAACGACCGCTATCAAAA  
 ATTTGATAACTCTATTGAAAACCAAATAAAGAATAAAAATATTA  
 CGTCAAAAAATAGAAAAATCTATCAAAAACATGGTATTACAAAAGA  
 GGTATTA AAAACTTACTATGCTAAAAAACAGTACGAGCTGACTGA

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FIG. 1 CONT'D

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MLHSKKIHSLSLIAVLSLATYTSLQPNHVAEQSQKTSTVLMSQKTIEHK  
LKVADKEAAPLYAKIDHIQRHIEVKKAKDLKVIELYINKDINQLEKQNK  
RLTKFYTSIDNQTWDSTSEVKKLIDKTTLSTNEKDRLKLYFEQRAYLET  
RLNDRYQKFDNSIENQNKELKILTSKIEKIYQKHGITKEVLKTYAKKTV  
RAD\*

ID-26

Clone 25

Clone 25 (partial sequence)

CTGAATTCCCAAAAACGCTACAATCAAACCTGGTATCCTACTTATGGTTTTTCTGAT  
ACTTATGCATTCATGGTACTAAAGAGTTTGCCAGACAGAATAAAAATCACCAAGAT  
CTCTGATCTCAAAAAGTTATCAACAACCTATGAAGGCAGGGGTTGATAGTTCATGGA  
TGAATCGCGAGGGAGATGGATACTGATTCGCTAAAACATACGGTTTTGAATTT  
TCACATATTTACCCTATGCAAATTGGCTTAGTCTATGATGCGGTTGAAAGTAACAA  
AATGCAATCTGTATTAGGCTACTCCACTGACGGTCGTATTCGAGCTATGATTTAG  
AAATTTTAAGGGATGATAAAAAATTCTTTCCTCCTTATGAAGCCTCTATGGTTGTCA  
ACAATTCTATCATCAAAAAGATCCTAAACTAAAAAATTACTCCATCGACTCGAT  
GGTAAAATCAATTTAAAAACGATGCAAAACCTTAATTATATGGTAGATGATAAACT  
TTAGAAGCTTGGCGTAATCATGGTCATAGCTGTTTCTGTGTGAAATTGTTATCCG  
CTCACAATTCCACACAACATACGAGCCGGAAGCATAA

LNSQKRYNQTYPTYGFSPTYAFMVTKEFARQNKITKISDLKKLSTTMKAGVDSSWM  
NREGDGYTDFAKTYGFEFSHIYPMQIGLVYDAVESNKMOSVLGYSTDGRISYDLEILR  
DDKKFFPPYEASMVVNNSIHKDPKLLKLLHRLDGKINLKTMQNLNYMVDKLEAW  
RNHGHSCFLCEIVIRSQFHTTYEPEA\*

ID-29

Clone 37

ATGAAAAAATTACTTTCCTAACATGTCTAATCATGATGTCTTTATGT  
TTAGTGGCATGTAAGCAAGCAATGTCGTCTAAGCAAGCAATGTC  
GTCTAAGCAAATTAAGATAAGAATAGTAAAGAAAAGGTGATTACT  
GTTGCAACTTACAGCAAACCTACATCTACCTTTTTAGATTTGATTAAA  
GATAATGTAAGAAAAGGATATACTTTAAAGGTTGTCATGGTCTC  
TGACTATATTCAGGCTAACATTGCTTTAGAAAACAAAGAACATGATG  
CTAACCTTTTACAACATGAATTTTTCATGAGTATCTTTAATAAGGAAA

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FIG. 1 CONT'D

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ATGATGGTCATCTAGTGTCAATTACACCAATTTATCATTTCATTGGCTG  
GTTTTTATGGTCAACATTTGAAAAATATTGCCGAGCTTAAAGACGGT  
GCTAAGGTAGCGATTCCGTCTGATCCTGCCAATATGACTAGAGCTCT  
GCTATTATTGCAAGAAAAGAACTTATCACCTTAAAGAATACGTCCA  
AAAAGACCAAGGCTATCGAAGATATTACTAACCCTAAAAAATTA  
CGAATTGAACCTGTAGCATTACTTAACCTCAATCAGGCCTATTTTGAA  
TATGACCTTGTCTTTAATTTCCCTGGATATGTGACAAAAATCAATCTA  
GTTCTAAAAGGGATAGATTATTATATGAGAAAAAACAGATATCCG  
TTTTGCAGGTGCCTTGGTAGCTCGTGAAGATAATAAAAATAGTGATA  
AAATAAAAGTACTTAAAGAAGTACTAACAAGTAAAGAGATTCGTCA  
CTATATCACTAAGGAGATTCCAAGTGAAGCAGACGTTGCGTTCTAG

MKLLSLTCLIMMSLCLVACTKQAMSSKQAMSSKQIKDKNSKEKVITV  
ATYSKPTSTFLDLIKDNVKEKGYTLKVVMSDYIQANIALENKEHDANL  
LQHEFFMSIFNKENDGHLVSITPIYHSLAGFYGQHLKNIAELKDGAKVAI  
PSDPANMTRALLLLQEKKLITLKNSTKTKAIEDIITNPKKLRIEVALLN  
LNQAYFEYDLVFNFPGYVTKINLVPKRDRLLYEKKPDIRFAGALVARED  
NKNSDKIKVLKEVLTSKEIRHYITKEIPSEADVAF\*

ID-30

Clone 38

CTGTTGGCTAAGGAAACCACTATGTCTGTCCTTTGGTATCAAATTCTGCAGAAGC  
CAAGGCTTTATATTTACAAGTTATAATGTTGCTAAAATGAAGTTAGATGATTGGT  
TACAAAAGCCCAGTGAAAAACCATATTCAATTATCTTAGATTTAGATGAAACAGTT  
TTAGATAATAGCCCATATCAAGCAAAGAATATTAAGATGGCTCTAGTTTCACGCC  
AGAGAGTTGGGATAAATGGGTGCAAAGAAATCAGCTAAGGCTGTTGCGGGTGCC  
AAAGAATTTTTGAAGTATGCTAATGAAAAGGGAATAAAAATTTATTATGTCTCAGA  
TCGTACAGATGCTCAAGTTGATGCGACTAAAGAAAATTTAGAGAAGGAAGGTATA  
CCTGTTCAAGGGAAAGACCACTTGCTTTTCCTTAAAAAAGGAATGAAATCTAAAGA  
GAGTCGCCGTCAGGCAGTTCAAAAAGATACCAATTTAATTATGCTTTTTGGAGATA  
ATTTAGTTGATTTTGCTGATTTTCTAAATCATCTAGTACAGATAGAGAACAACACTAC  
TAACTAACTTCAAAGTGAGTTTGGTAGTAAATTTATTGTTTTCCCAAATCCTATGT  
ACGGTTCTTGGGAAAGTGCTATTTATCAAGGAAAACATCTGGATGTTCAAAAACAA  
TTGAAAGAACGACAAAAAATGTTGCATTTCGTATGATTAA

MAKLTVKDVDLKGKKVLRVDFNVPLKDGVITNDNRITAALPTIKYIIEQGGRILFSH  
LGRVKEEADKEGKSLAPVAADLAAKLGQDVVFPVTRGAKLEEAINALEDGQVLLVE  
NTRFEDVDGKKESKNDEELGKYWASLGDGIFVNDAFGTAHRAHASNVGISSNVEKAV  
AGFLENEIAIYIQEAVETPERPFVAILGGSKVSDKIGVIENLLEKADKVLIGGGMTYTFY  
KAQGIEIGTYLEKEDKLDVAKDSZ

ID-31

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FIG. 1 CONT'D

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Clone 41

ATGGATAATAAAGGTAATAACGCCAATGTGATTGATGCAATCGCTGAGGGTGCAA  
GCACAGGTGCACAAATGGCTTTCTCAATTGGTGCTAGTTTGATTGCCTTTGTTGGTT  
TAGTTTCTTTGATTAA

MDNKGNNANVIDAIAEGASTGAQMAFSIGASLIAFVGLVSLI

ID-32

Clone 42

ATGAAAAAGAAAAACAAATCCTCTAACATTGCTATAATTGCAATCTT  
TTTTGCTATTATGCTTGTCATTCATTTTTTGTCATCATTTATTTTTAGTT  
TTTGGTTAGTCCCTATTAACCTACTTTGATGCATATCCCAGTTATTA  
TTGCATCTATAGCCTATGGACCTCGTATTGGTGCAACTCTAGGCGCCT  
TAATGGGGGGGATCAGCGTAGCTAACAGCAGCATTGTTCTATTACCA  
ACGAGTTACCTCTTCTCACCTTTTGTTGAAAATGGTAATTTTTATTTCG  
CTAATTATTGCACTTGTACCACGTATTCTAATCGGGATTATTCCTTAT  
TTCGTTTACAAATTAATAACAACCGCTTTGGTTTGGCTATCTCAGGT  
GCTATAGGCTCTCTAACAACACAGTATTTGTTTTATCTGGAATTTTT  
ATCTTTTTTTCAAGTACTTATAATGGGAATATCAAGCTAATGCTCGCT  
GGGATTATTTTCACTAATTCATTAGCTGAGATGGTCATTGCAGCTATC  
ATTGTATATCTAACTGATCCTCGTATTCTCAATATTAACATTAA

MKKKNKSSNIAIIAIIFFAIIMLVIHFLSSFIFSFVLPVPIKPTLMHIPVIIASIA  
Y GPRIGATLGALMGGISVANSSIVLLPTSYLFSFPVENGNFYSLIIALVPRILI  
GIIPYFVYKLLHNRFLAISGAIGSLTNTVFVLSGIFIFSSTYNGNIKLML  
AGIISSNSLAEMVIAAIIIVYLTDPRIKLNKH\*

ID-33

Clone 43

TTGAATATGACATTACAAGACGAAATCAAAAAACGCCGTACTTTTGCCATCATCTC  
TCACCCGGATGCTGGTAAGACGACTATTACTGAGCAATTATTATTTTTGGTGGTG  
AAATTAGAGAAGCAGGGACAGTAAAAGGGAAAAAATCAGGTACTTTTGCAAAGTC  
CGACTGGATGGATATTGAAAAGCAACGGGGTATCTCTGTTACTTCATCTGTTATGC  
AATTTGATTACGCGGGTAAACGTGTTAA

MNMTLQDEIKKRRTFAIISHPDAGKTTITEQLLYFGGEIREAGTVKGKSGTFAKSDW  
MDIEKQRGISVTSSVMQFDYAG

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FIG. 1 CONT'D

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KRV

ID-34

Clone 44

ATGGCAGATAAAAACAGAACATTTAAACTTGTAGGTGCAGGATCTTC  
TAGCACACAAGAAAAAATTGAAAAGCCTGCTCTTTCGTTTATGCAAG  
ATGCGTGGCGTCGCTTGAAAAAACAATTAGCAGTAGTTTCACTC  
TATTTATTAGCTCTTTACTTACTTTTTTCGTTAGCCTCAAATTTATTG  
TAACTCAGAAGGATGCTAATGGGTTTGATTTCGAAAAAAGTAACGACA  
TATCGCAACTTACCACCTAAATTGAGTTCAAACCTTCCTTTTTGGAAT  
GGTAGCATTAATCCATCA

MADKNRTFKLVGAGSSSTQEKIEKPALSFMQDAWRRLKKNKLA VVSLY  
LLALLLTFSLASNLVFTQKDANGFDSKKVTTYRNLPPKLSSNLPFWNGSI  
NPS

ID-35

Clone 46

ATGAAAAGAAAACAGTTTATAAAATTAGGAATTGCAACCTTACTAACGGTTATTT  
GCTTTACACACCAATAAACCTAGCTACAAATCATAACCACAGAAAATATTGTTACTG  
CTCAAGAGTATAAAACAAAGAGAATGGTACTTTACCTTTTAA

MKRKQFIKLG IATLLTVISLYTPINLATNHTTENIVTAQEYKTKENILFLL

ID-36

Clone 50

ATGTTTTATAATCCTTTACTTTTTATTGTTACTAATTACAATTGCTGTATTTTTCTTAG  
CTAAGAAAAAATGGCAATTACCGACATTTACTTTCATTGGTTTGCTATTTATCTATA  
ACCAAGGGCTGTGGGAACAGTTGATTAAT

MFYNPLLFIVLITIAVFFLAKKKWQLPTFTFIGLLFIYNQGLWEQLIN

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FIG. 1 CONT'D

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ID-37

Clone 51/52

GTGGTGCAAATAATGAAAAACATATAAAAAGTATCATACCAATAGT  
TCTTATTGGTATGATACTAGGAGGCTGTCAAATGAATAGTGAACATA  
AAAGTCAGTATAATGAAACAAAAAGTAGCAAGCAATCAGAAGTGAA  
GAAAGATAAAAAAATGACAAAAAAGAACAATTAGCTTATCTCAA  
GAGCATGAACAAGAAATAATTGATTTTGTAAAATCTCAGAATAAAAA  
GATAGAATCTGTACAAATTGATTGGAATGATGTTTCGATGGAGTAAAG  
GGGGAAATGGTACACCTCAAGGAGGAGGAGAGGGGATTTTACTTTT  
GGGGAGATTAATAATGATTCTGAATCAAGTTGGAGAGTTGATATTGA  
TATAGAAAAAGGACGGCTAGACCTAAAAAATATGTATTTAGGACAA  
CCTATACGAATTGGAGGTAATTATTTGAGTAA

MVQIMKKHIKSIPIVLIGMILGGCQMNSEHKSQYNETKSSKQSEVKKDK  
KMTKKEQLAYLKEHEQEIIDFVKSQNKKIESVQIDWNDVRWSKGGNGT  
PQGGGEGILLFGEINNDSESSWRVDIDIEKGRLLDLKNMYLGQPIRIGGKLF  
E\*

ID-38

Clone 53

ATGGAATTTTTGGCTTATAATGCTTTCACAGCAATCGGTGTTTCTATT  
CCGCACGGTAATCATTTCCTTTTACTATAAAGGATATGTCTCCA  
TTAGAGTTAGAAGCAACAAGGATGGTGGCAGAGCATAGAGGACATC  
ATATTGATGCATTAGGGAAAAAAGATTCTACAGAGAAACCAAAGCA  
TATTTCTCATGAACCTAATAAGGAACCTCACACAGAGGAAGAACC  
ATGCAGTAACACCGAAAGACCAACGTAAAGGCAAACCAAATAGCCA  
GATTGTCTACAGTGCTCAAGAAATTGAAGAGGCAAAAAAAGCTGGT  
AAATACACAACATCTGATGGTTACATTTTTGATGCTAAAGATATTAA  
AAAAGATACAGGTACAGGTTATGTCATTCCACATATGACACATGAGC  
ATTGGGTACCAAAGAAAGATTTATCAGAGTCGGAATTAAGCAGCT  
CAAGAATTTCTTTCAGGAAAATCTGAAGCAAATCAAGACAAACAAA  
AACAGGTAAAACAGCTCAAGAAATCTATGAGGCAATTGAACCAAAA  
GCAATTGTTAAACCTGAAGATTTATTATTTGGAATTGCACAAGCGAC  
AGACTATAAGAATGGTACATTTGTAATTCCTCATAAAGATCATTACC  
ATTATGTGGAATTAAGATGGTTTGTGATGAAGAAAAAGATCTTTTAGCT  
GATTCAGATAAGACATATTCTTTAGAAGACTATTTAGCTACGGCTAA  
ATATTACATGATGCACCCAGAAAAACGTCCTAAAGTTGAAGGATGGG  
GTAAAGATGCTGAAATTTATAAGGAAAAGGACTCTAATAAAGCAGA  
TAAACCAAGTCCTGCACCAACTGATAATAAATCAACATCAAATTCTA

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FIG. 1 CONT'D

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GTGACAAAACTTAAGTGCTGCAGAAGTATTCAAACAAGCAAAACC  
 AGAAAAAATTGTACCGCTTGATAAAATTGCTGCTCACATGGCATATG  
 CAGTTGGATTTGAAGATGATCAATTGATTGTTCCCTCATCATGATCATT  
 ATCATAATGTTCCATATGGCATGGTTTGACAAGGGTGGTTTTATGGAAA  
 GCACCAGAAGGCTATACATTACAACAACCTCTTCTCAACAATTAATA  
 CTACATGGAACATCCTAATGAATTACCAAAAGAAAAGGGTTGGGGA  
 CACGACAGTGATCATAACAAAGGCTCAAATAAAGACAATAAAGCCA  
 AAAATTATGCTCCAGATGAAGAACCTGAAGATTCAGGGAAAGTAACT  
 CACAACATATGGTTTTTATGATGTTAATAAAGGTTTCAGACGAAGAAGA  
 ACCAGAAAAACAAGAAGATGAATCAGAGCTAGATGAATATGAACTA  
 GGAATGGCACAAAACGCTAAGAAATATGGTATGGATAGACAATCTTT  
 TGAAAAGCAACTCATCCAATTATCAAATAAATATAGTGTAAGTTTTG  
 AAAGC

MEFLAYNAFTAIGVSIPHGNHFHFIHYKDMSPLELEATRMVAEHRGHHI  
 DALGKKDSTEKPKHISHEPNKEPHTEEHHA VTPKDQRKGKPN SQIVYS  
 AQEIEEAKKAGKYTTSDGYIFDAKDIKDTGTGYVIPHM THEHWVPKK  
 DLSESELKAAQEFLSGKSEANQDKPKTGKTAQEIEAIEPKAIVKPEDLL  
 FGIAQATDYKNGTFVIPHKDHYHYVELKWFDEEKDLLADSDKTY SLED  
 YLATAKYMMHPEKRPKVEGWGKDAEIKYKEKDSNKADKPSAPT DNK  
 STSNSSDKNLSAAEVFKQAKPEKIVPLDKIAAHMAYAVGFEDDQLIVPH  
 HDHYHNVPMAWFDKGGLWKAPEGYTLQQLFSTIKYYMEHPNELPKEK  
 GWGHDS DHNKGSNKDNKAKNYAPDEEPEDSGKVTHNYGFYDVNKGS  
 DEEPEKQEDESELDEYELGMAQNAKKYGM DRQSF EKQLIQLSNKY SV  
 SFES

ID-39 (Same as ID-76)

Clone 56

ATGAGGAAACGTTTTTCCTTGCTAAATTTTATTGTTGTTACTTTTTATT  
 TCTTTTTCTTTATTCTTTTTCCGCTTTTTAAGGCCAAAGATTGTCAGGT  
 TGTTTTATGCAAGTTTTCAAGGAGATCATTGGGACATTTGTAACGCATT  
 TGATTTTCCGTATTTACATCGCTTTGATCTCATTAAAGGTAAAGAAAA  
 TCAACTTTACTTTATAGGTTGTACAATTGCTAACAGTAAAGCCTACAC  
 TGAGGATTGGAGTGATAAAGGCCGAATTTTTGTTGCTCGTTTTAATAC  
 TCAAACCATACATTGGAAGGATTGCAACAATTGCCTCAAAC TTTAT  
 TAAAAAATCATGGATACTATGCCATTCAGGATGAAGGATATTCATTG  
 ATTACTTCAGTAGAAGGGGTACTCAAAC TCACTTATCCAGAATTTTCT  
 ACTACAGGCGACTGGCAATTAGAACGGCTTTTCGATGAGGAGACAAG  
 CGATGTGGTGAAAGTGGATATTAATCAGGATGGTAAGGATGAGTATG  
 TGATCATCCAAGGTTTTCATGGAGATCGTTTACGTATCTTCACTGAAG  
 ATTTCCGGTCGAGAATTATCCATTATCCTGAAAAAACCCCATTTGGTC  
 ACGCTATTTGGAGTGGTCGTTTACTTAATCAGACTTGTTTCGTATT CG

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FIG. 1 CONT'D

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GGTGGCGATCAGAAAAAGCAGAATTAAGGCTTTTTCACTTTGTAGAT  
 GGGCACTTGGTTTCAGAATTAGTAGATGCAAAAAGCAGCTTCTAGTAA  
 TGTCTTAGCTTTTGAAAAAGATGGAAAAGCTTATCTTTTCTCAGCCAA  
 TAACGGACGTGGCGAAGTTGCTCTTTATCAATTAGTAAAATAA

MRKRFSLLNFIVVTFIFFFIFLPLFKAKDCQVVYASFQGDHWDICNAFDF  
 PYLHRFDLIKGENQLYFIGCTIANSKAYTEDWSDKGRIFVARFNTQNHT  
 LEGLQQLPQTLLKNHGYIAIQDEGYSLITSVEGVCLKLTYPEFSTTGDWQ  
 LERLFDEETSDVVKVDINQDGKDEYVIIQGFHGDRLRIFTEDFGRELFHY  
 PEKTPFGHAIWSGRLLNQTCFVFGWRSEKAELRLFHFVDGHLVSELVDA  
 KAASSNVLAFEKDGKAYLFSANNGRGEVALYQLVK\*  
 ID-40

Clone 57

ATGAAGCACAAGTTAAAAGCTTTTACGCTTGCTTTACTCTCAATATTC  
 TTTGTGTTTGGTGGAAAGGTCAGTGCAGAGACTGTGAATATTGTTTCT  
 GATACAGCATAACGCTCCATTCGAATTTAAAGATTCTGATCAAACCTTAT  
 AAAGGAATCGATGTTGACATCGTTAACGAAGTCGCTAAGCGTGCTGG  
 CTGGAATGTTAACATGACGTATCCAGGTTTTGATGCCGCAGTTAACG  
 CTGTTCAATCTGGACAGGCAGATGCGCTAATGGCCGGAACACTGTT  
 ACTGAAGCACGTAAAAAAGTCTTTAATTTCTCAGATACTTATTACGAT  
 ACTTCCGTTATTCTTTATACTAAAAATAATAATAAAGTCACAAACTAC  
 AAACAACATAAAGGAAAAGTAGTCGGTGTAATAAATGGAACAGCTG  
 CTCAAAGCTTCTTAGAAGAAAATAAATCTAAATACGGCTATAAAGTT  
 AAAACATTTGATACAAGCGACCTAATGAATAACAGCCTTGATTCTGG  
 TTCTATTTACGCCGCTATGGACGATCAACCAGTTGTGCAATTTGCGAT  
 AAATCAAGGAAAAGCTTACGCCATTAACATGGAAGGCGAAGCAGTT  
 GGTAGCTTTGCATTTGCTGTCAAAAAAGGTAGTGGACACGATAATCT  
 AATTAAGAATTTAACACAGCTTTTGCACAAATGAAATCAGATGGCA  
 CTTATAATGACATCATGGATAAATGGCTTGGAAAAGACGCTACAAAA  
 ACAAGCGGCAAAGCAACAGGTAATGCCAATGAAAAGCAACTCCTG  
 TAAAGCCAAGTTATAAAATTGTTTCTGATTCTTCATTCGCACCATTCTG  
 AATATCAAAACGGTAAAGGGAAATATACTGGTTTTGATATGGAATTA  
 ATCACGAAAATTGCTAAACAGCAAGGTTTTAACTTGATATCTCAA  
 TCCAGTTTTGATGCCGCTTTAAATGCTGTCCAATCTGGGCAAGCTGA  
 CGGTGTTATTGCAGGAGCCACAATCACAGAAGCACGCCAAAAAATCT  
 TTGATTTTTCTGATCCTTATTACACATCTAGCGTTATCTTAGCGGTTAA  
 AAAAGGAAGCAATGTCAAATCATACCAAGATTTAAAAGGAAAAACA  
 GTTGGTGCTAAAAATGGTACTGCCTCATATACTTGGTTATCAGACCAC  
 GCAGATAAGTACAACATCATGTTAAAGCATTGATGAAGCATCTAC  
 AATGTATGATAGTATGAACTCAGGTTCAATTGATGCTCTAATGGATG  
 ACGAAGCCGTTCTTGCTTACGCTATTAATCAAGGTCGTAATTTGAA  
 ACACCTATCAAAGGTGAAAATCAGGCGATATCGGATTTGCAGTGAA

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FIG. 1 CONT'D



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AAAAGGGGCAAATCCAGAATTAATTAATAATGTTTAAACAACGGTCTTG  
 CTTCACTCAAAAAATCGGGTGAGTACGATAAACTTGTTAAAAAATAC  
 CTTTCCACAGCCAGCACTTCTTCAAACGATAAAGCTGCTAAACCTGT  
 AGATGAATCAACTATTTTAGGGTTAATTTCTAATAACTACAAACAATT  
 GCTATCTGGTATTGGAACACTTTAAGTTTAACTCTTATCTCGTTTGC  
 GATTGCTATGGTTATTGGTATTATCTTTGGTATGATGAGCGTATCACC  
 AAGTAATACTCTCCGCACAATTTCAATGATTTTTGTTGATATTGTCCG  
 TGGTATTCCACTCATGATTGTGGCCGCTTTATTTTCTGGGGTATTCTT  
 AATTTAATCGAAAGCATCACAGGTCACCAAAGTCCAATTAATGACTT  
 CGTTGCTGCTACTATCGCTCTTTCTTTAAATGGTGGTGCGTACATTGC  
 TGAAATTGTACGTGGTGGTATTGAAGCTGTTCTTCTGGTCAAATGGA  
 AGCAAGTCGCAGCTTAGGTATTTCTTACGGCAAACTATGCAAAAGG  
 TTATCTTACCTCAAGCAGTACGCCTTATGTTACCAAACCTTTATCAACC  
 AATTTGTCATCTCATTAAAGGATACAACAATTGTATCAGCAATCGGA  
 CTTGTGGAACCTTCCAAACTGGTAAATCATAA

MKHKLKAFTLALLSIFVFGGKVS AETVNI VSDTAYAPFEFKDSDQTYK  
 GIDVDIVNEVAKRAGWNVNMTYPGFDAAVNAVQSGQADALMAGTTV  
 TEARKKVFNFSDTY YDTSVILYTKNNNKVTNYKQLKGKVVGVKNGTA  
 AQSFLEENKSKYGYKVKT FDTSDLMNNSLD SSGSIYAAMDDQPVVQFAI  
 NQGKAYAINMEGEAVGSFAFAVKKGSGHDNLIKEFN TAF AQMKSDGTY  
 NDIMDKWLGKDATKTS GKATGNANEKATPVKPSYKIVSDSS FAPFEYQ  
 NGKGKYTGFDME LITKIAKQQGFKL DISNPGFDAALNAVQSGQADGVIA  
 GATITEARQKIFDFSDPYTSSVILAVKKGSNVKS YQDLKGKTVGAKNG  
 TASYTWLSDHADKYN YHVKA FDEASTMYDSMNSGSIDALMDDEAVLA  
 YAINQGRKFETPIKGEKSGDIGFAVKKGANPELIKMFNNGLASLKKSGEY  
 DKLVKKYLSTASTSSNDKAAKPVDESTILGLISNNYKQLLSGIGTTLSLTL  
 ISFAIAMVIGIIFGMMSVSPSNTLR TISMIFVDIVRGIPLMIVA AFIFWGIPN  
 LIESITGHQSPINDFVAATIALSLNGGAYIAEIVRGGIEAVPSGQMEASRSL  
 GISYGKTMQKVILPQAVRLMLPNFINQFVISLKDTTIVSAIGLVELFQTGK  
 S\*

ID-41

Clone 58

TTGGAAGGTTTACTTATTGCATTGATTCCCATGTTTGCGTGGGGAAGTATTGGATT  
 GTTAGTAATAAAATTGGAGGGCGTCCAAATCAACAAACATTTGGAATGACTTTAGG  
 AGCATTGCTATTTGCGATTATCGTATGTTTATTAA

MEGLLIALIPMFAWGSIGFVSNKIGGRPNQQTFGMTLGALLFAIIVCLF

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FIG. 1 CONT'D

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 ID-42

Clone 70

ATGAATACTATTTATAATACATTGAGAACAGATAAAAGGTTATAAAGT  
 TTATGAGGGGTATTTATATGAAATTACTGGTGAAGAATGTGAAGAAG  
 CCTTAGACCTTGTGATTCCTAAGAATATTGTATTTGCAGATACAGATA  
 CTTGTGGCTACACTTTTTTACTCAATGAAGATGGAACAGTTTATGATG  
 ATGTGACTTTCTACAAATTTGATGATAAATATTGGTTGGCTAGTCATA  
 AAGCTTTGGATTCTTATTTAGACAACATCAATTTTACTATAACCGTAA  
 CAGATATTTCTGACGAGTATAAAATGCTGCAAATTGAAGGAAGATAT  
 TCGGGAGAAATTGCTCAGTCATTTTATGAATATGATATTTCAACACTT  
 AATTTTCGTACTCTTCGCATAGAGATGGACTTCATCAAAGGTGAGGA  
 AAGGTTATCTTGGCGTAGATTTGGTTTTTCTGGAGAATTTGGCTATCA  
 ATTTTTCCTACCATCTTCTATTTTGTACTTTTGTTCGGATGTCTGT  
 GAAGGTATAGCAGAGTGTGGGGATGAACTTGATAGATATTTAAGGTT  
 TGAAGTGGGACAACCCATTACTGATATTTATCAACAAGAAGAATATT  
 CTTTATATGAAATAGGTTATTCTTGGAAATCTAGATTTACAAAGGAA  
 GAATTTAGAGGTTCGCGATAGCTTGTTAGAGCACATCAGATCAGCAAC  
 AGTTAAAAGTGTTGGATTCTCAACGAAGGAAAAACTCGCTTCAGGAA  
 CACCAGTGCTATTTGATGACCAAATTGTTGGAAAGATTTTTTGGATAG  
 CAGACGAGAAACACTCTTCGGAAAATTACCTAGGTTTGATGATTGTT  
 AACCAAACATATGCTCATTGAGGAGTTACTTTTGTAACAGAAGATGG  
 CCAAATTTTGAAAACACAATCAAGCCCTTATTGTATCCCAGAAAGTT  
 GGAACAAAGAATGA

MNTIYNLRLTDKGYKVYEGYLYEITGEECEEALDLVIPKNIVFADTDTCG  
 YFLLNEDGTVYDDVTFYKFDDKYWLASHKALDSYLDNINFDYTVTDIS  
 DEYKMLQIEGRYSGEIAQSFYEYDISTLNFRTLRIEMDFIKGEERLSWRRF  
 GFSGEFGYQFFLPSSIFATFVSDVCEGIAECGDELDRYLRFEVGQPITDIY  
 QQEESLYEIGYSWNLDFTKEEFRGRDSLLEHIRSATVKS VGFSTKEKLA  
 SGTPVLFDQIVGKIFWIADEKHSSENYLGMLMIVNQTYAHSVTFVTE  
 GQILKTQSSPYCIPESWNKE\*

ID-43

Clone 78/94

ATGGAGTTAGTAATTAGAGATATTCGTAAGCGGTTTCAGGAAACAGA  
 GGTCTTGAGAGGAGCAAGTTACCGATTTTATTCAGGTAATAACAG  
 GGGTCTTAGGTAGGAATGGTGCTGGGAAAACAACCTTTATTTAATATA  
 CTTTATGGGGATCTTGCAGCTGACAACGGGACCATTTGTTTATTGAAG  
 GATAATCACGAGTATCCTCTTACCGATAAAGGATATTGGTATTGTTTAT

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FIG. 1 CONT'D

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TCCGAAAACCTACCTTCCAGAATTTTAAACAGGGTATGAATTTGTAAA  
ATTTTACATGGATTTACATCCTTCAGATGATTTAATGACAATAGATGA  
TTATTTAGATTTTATGGAAATAGGACAAACAGAGCGTCATAGAATTA  
TCAAAGGATATTCTGATGGAATGAAGAGTAAGCTCTCATTAAATTTGC  
CTGATGATTTCTAAGCCAAAAGTAATTTTACTAGATGAGCCACTGAC  
TGCAGTTGATGTTGTATCAAGTATTGCAATAAAACGCCTTTTGTGGGA  
ATTAAGTGAGGATCATATTATTATATTATCAACTCATATAATGGCCTT  
AGCAGAAGATCTATGTGATATTGTGGCTGTATTAGACAAAGGAAAAC  
TCCAAACATTAGATATTGATCGTAAACATGAACAATTCGAAGAGCGT  
CTTCTTCAAGTGTTGAAGGGAGATGAATATGACAAGTAA

MELVIRDIRKRFQETEVLRGASYRFYSGKITGVLGRNGAGKTTLFNILYG  
DLAADNGTICLLKDNHEYPLTDKDIGIVYSENYLPEFLTGYEFVKFYMD  
LHPSDDLMTIDDYLDMEIGQTERHRIKGYSDGMKSKLSLICLMISKPK  
VILLDEPLTAVDVVSSIAIKRLLLELSEDHIIILSTHIMALAEDLCDIVAVL  
DKGKLQTLDIRKHEQFEERLLQVLKGDEYDK\*

ID-44

Clone 80

TTGTTTATGAGATATACAAATGGAAATTTTGAAGCCTTTGCAAGACCT  
CGAAAACCTGAAGGTGTGGATAAAAAATCCGCTTATATTGTTGGTTC  
TGGTTTAGCAGGATTAGCTGCCGCTGTCTTTTTAATACGTGACGGTCA  
AATGGATGGTCAACGTATTCATATTTTGAAGAACTACCTCTTTCTGG  
AGGATCACTTGACGGTGTCAAACGACCTGATATCGGTTTTGTAAACGC  
GTGGTGGTCGTGAAATGGAAAATCACTTCGAATGTATGTGGGATATG  
TACCGTTCCATCCCCTCTCTCGAAGTTCAGATGCTTCTTATCTAGAT  
GAATTTTATTGGCTTGACAAGGATGATCCCAATTCATCTAACTGTGCG  
CTCATTATAAACAGGGGAATCGCTTAGAATCTGATGGTGATTTTAC  
ACTCGGAACACATTCCAAAGAGTTAGTTAAGCTAGTCATGGAGACTG  
AAGAGTCTTTAGGTGCTAAGACGATTGAAGAAGTTTTTTCAAAGAA  
TTTTTTGAAAGTAATTTTGGACTTATTGGGCTACTATGTTTGCCTTTG  
AGAAATGGCATTACAGCGATTGAAATGCGTCGATATGCTATGCGCTTT  
ATCCATCATATTGGTGGTCTGCCTGATTTCACTTCATTAATAATTAAT  
AAATATAATCAATATGATTCTATGGTGAAACCAATCATCAGTTATTTA  
GAGTCTCATAATGTAGATGTTCAATTTGATAGCAAGGTAACATAAT  
CTCCGTAGACTTT

MFMRYTNGNFEAFARPRKPEGVDKKSAYIVGSLAGLAAAVFLIRDGQ  
MDGQRIHIFEELPLSGGSLDGVKRPDIGFVTRGGREMFECMWDMY  
RSIPSLEVPDASYLDEFYWLDKDDPNSSNCRLIHKQGNRLES DGDFTLGT  
HSKELVKLVMETEESLGAKTIEEVFSKEFFESNFWTYWATMFAFEKWHS

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FIG. 1 CONT'D

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AIEMRRYAMRFIHHIGGLPDFFTSLKFNKYNQYDSMVKPIISYLESHNVDV  
QFDSKVTNISVDF

ID-45

Clone 81

TTGTTGGCTTCTTTATTTATCGTCCGTTTGTCAAAAATCGCTTTCGCTAA  
GGAGGAGCAATATGAAAAAATTACTTAGATGGCTTCCTCCTGTACTT  
TTCATTATTATCCTTATAGGAATGACTATCTTAGGTAAGTCCTATATC  
AATAAAGTAACAGCTCACAAAATAAACTCTATAACTCTCGAATGAC  
TCCTACTATTTTAATTTTCAGGATCCAGTGCTACTCAAGAACGATTTAA  
CAGCATGTTAGCACAGCTCAACCAAATGGGAGAAAAACATAGCGTTT  
TAAAGTTAACTGTCAAAAAAGACAATAGCATTATCTACAATGGACAA  
ATTAGCGGCAATGACCACAAACCCTACATTGTCATTGGATTTGAAAA  
TAATGAAGATGGTTATAGTAACATCAAAAAACAAACAAAATGGCTA  
CAGATTGCTATGAATGATCTTCAGAAGAAATATAAATTTAAACGTTT  
TAACGCTATCGGTCATTCAAATGGTGGCTTATCATGGACTATTTTCCT  
AGAAGATTATTACGACTCTGATGAATTTGATATGAAATCATTGTAA  
CAATGGGAACACCTTTTAACTTTGAAGAAAGTAACACCTCAAATCAT  
ACTCAAATGCTTAAAGATTTAATCAGTAATAAAGGAAATATTCCATC  
AAGTCTCATGGTATACAATTTGGCAGGAACTAATTCATATGATGGTG  
ATAAAATTGTTCCATTTGCTAGTGTGGAGACTGGTAAATATATTTTCC  
AAGAAACCGCTAAACACTATACCCAATAACAGTAACTGGTAATAAT  
GCTACACATTCTGACTTGCCTGATAATCCTGAAGTTATCCAATATGTC  
GCAGAAAAAATTCTTAAAAATGAGAAAGGTAAATTACCAAACCTC  
ACTAA

MLASLFIVRLSKSLRRSNMKKLLRWLPPVLFIIILIGMTILGKSYINKVT  
AHKIKLYNSRMTPTILISGSSATQERFNSMLAQLNQMGEEKHSVLKLTVK  
KDNSIIYNGQISGNDHKPYIVIGFENNEDGYSNIKKQTKWLQIAMNDLQK  
KYKFKRFNAIGHNSNGGLSWTIFLEDYYSDEFDMKSLLTMGTPFNFEES  
NTSNHTQMLKDLISNKGNISSLMVYNLAGTNSYDGDKIVPFASVETGK  
YIFQETAKHYTQLTGTGNNATHSDLPDNPEVIQYVAEKILKNEKGKLPK  
PH

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ID-46

Clone 83

TTGAAATTAGGTATTACAACATTCGGAGAGACAACAATCCTTGAAGAAACAAACC  
AAAGCTATTCACATCCTGAGAGGATTCGCCAATTAGTTGCTGAGATTGAACTAGCT  
GATCAAGTTGGTTTAGATGTATATGGTATTGGAGAGCACCATCGTGAAGATTTGC

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FIG. 1 CONT'D

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GGTCTCTGCACCCGAAATTATCCTAGCAGCAGGAGCGGTTAGAACTAATAATATCC  
GTTTATCTAGTGCAGTAACGATTCTCTCTTCCAATGATCCTATTCGCGTCTATCAGC  
AATTTTCAACGATTGACGCACTTTCAAATGGTAGAGCAGAAATTATGGCAGGGCGT  
GGTTCCTTTATTGAGTCTTTTCCATTGTTTGGATACGATTTAGCGGATTATGATGAT  
TTATTTAATGAAAAAATGGATATGTTGTTAGCAATTAAGTTCAGCGACAAATCTCGA  
TTGGAAGGTCATTTGACACAAACAGTTAATGAGCGACCAATTTATCCAAGAGCAT  
TACAAAGACAGTTATCAATATGGGTGGCAACAGGAGGAAATGTTGATTCTACAATT  
CGTATTGCAGAACAAGGTTTGCCAATTGTTTATGCAACTATTGGTGGGAATCCCAA  
AGCCTTTCGTCAATTGGTCCATATTTATAAAGAAGTTGGTAAGTCCGTAATGGACA  
CAAACCAGGAACAATAAAAGTTGCTGCTCACTCTTGGGGATGGATTGAAGAGGA  
TAATCAAACCGCTATTGACCGTTATTTTTTCCCTACGAAACAGACCGTCGATAATAT  
TGCTAAGGGACGCCCTCATTGGTCTGAAATGACTAAAGAGCAGTATTTACGTTCAA  
TAGGTCCAGAAGGTGCTATTTTTGTAGGAAATCCTGAAGTGGTTGCACATAAAATT  
ATAGGACTTTGGTGA

MKLGITTFGETTILEETNQS YSHPERIRQLVAEIELADQVGLDVY GIGEHHRDFAVSAP  
EIIAAGAVRTNIRLSSAVTILSSNDPIRVYQQFSTIDALSNGRAEIMAGRGSFIESFPLF  
GYDLADYDDL FNEKMDMLLAINSATNLDWKGHLTQTVNERPIYPRALQRQLSIWVAT  
GGNV DSTIRIAEQGLPIVYATIGGNPKAFRQLVHIYKEVGKSVMDTNQEQLKVAHSW  
GWIEEDNQTAIDRYFFPTKQTVDNIAKGRPHWSEMTKEQYLR SIGPEGAIFVGNPEVV  
AHKIIIGLW

ID-47

Clone 86

ATGATAGAGTGGATTCAAACACATTTACCAAATGTATATCAAATGGG  
TTGGGAAGGTGCTTACGGCTGGCAGACAGCTATTGTACAAACCCTTT  
ATATGACTTTTTGGTCGTTCCCTTATTGGAGGTTTAATGGGATTGTTAG  
GAGGTTTATTCCTTGTTTTAACTAGTCCTAGAGGAGTTATTGCTAATA  
AATTAGTATTTGGAGTTTTAGATAAAGTTGTTTCTGTTTTTAGAGCTC  
TGCCCTTCATTATTCTTCTTGCTTTGATTGCGCCAGTAACTCGCGTAAT  
TGTAGGAACAACACTTGGTTCACCAGCAGCTTTGGTACCTCTTTCTTT  
GGCAGTTTTCCCATTTTTTGCTCGTCAAGTTCAAGTTGTTTTAGCTGA  
ACTTGATGGTGGAGTTATTGAGGCTGCACAAGCCTCAGGTGGAACAC  
TTTGGGATATTATTGTAGTTTATCTTCGTGAAGGTCTACCAGATTTAA  
TTCGAGTATCAACGGTTACTTTGATTTCTTTAGTAGGTGAAACAGCTA  
TGGCTGGCGCTATTGGTGCAGGAGGATTGGGTTCTGTTGCTATTACTA  
AAGGATATAACTATTCTCGTGATGATATTACTTTAGTAGCGACTATTC  
TGATTTTTATTATTAATTTTCTTTATCCAATTTTTAGGTGATTTTTTAA  
ACGTCGCTTGAGTCATAAATAA

MIEWIQTHLPNVYQMGWEGAYGWQTAIVQTLYMTFWSFLIGGLMGLL  
GGLFLVLTSPRGVIANKLVFGVLDKVVSVFRALPFIILLALIAPVTRVIVG

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FIG. 1 CONT'D

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TTLGSPAALVPLSLAVFPFFARQVQVVLAEVDGGVIEAAQASGGTLWDII  
 VVYLREGLPDLIRVSTVTLISLVGETAMAGAIGAGGLGSVAITKGYNYSR  
 DDITLVATILILLIFFIQFLGDFLTRRLSHK\*

ID-48 (same as ID-43)

ID-49

Clone 96

TTGGCAGTTAGTTTTTCATGAAGTATTTGGTTGGGATTCTGCTTTTTTTA  
 TTATGATTATCAATATTCCATTGCTCCTTCTTTGCTACTTTGGCTTAGG  
 TAAACAAACCTTTTTAAAACTGTCTATGGTTCTTGGATTTTTCTGT  
 TTTTATTAAGTTAACACAAAGTGTACCAACTTTGACCCACAACCTCACT  
 CCTCGCAGCACTTTTTGGAGGTGTTATTGTAGGATGTGGTTTGGGGAT  
 TGTTTTTTGGAGCGACTCTTCAACTGGTGGAACGGGGATTATCATTCA  
 ATTCTTAGGAAAATATACTCCTATAAGCCTTGGACAAGGGGTTATAT  
 TGATTGATGGACTTGTTACAATTGTTGGTTTCTAGCTTTTGACAGTG  
 ATACGGTTATGTTTTCTATTATTGGGTTGATAACTATTAGTTATATTAT  
 TAATGCTATCCAAACTGGATTTACAACCTTAAGCACTGTCTTAATCGT  
 TTCTCAAGAGCACCAAAAAATTAAGACATATATCAATACTGTCGCAG  
 ATAGAGGAGTAACAGAAATTCCCGTTAAAGGGGGATATTCTGGAAC  
 AATCAAATCATGCTTATGACAACCTATTGCTGGTTATGAGTTTGCTAAA  
 TTACAAGAGGCAATAGCAGAAATTGACGAAACAGCCTTCATAACAGT  
 AACTCCAACATCACAAGCTTCTGGACGTGGATTTAGTCTTCAAAAAA  
 ATCATGGACGTCTTGATGAAGACATTCTTATGCCAATGTAA

MAVSFHEVFGWDSAFFIMIINIPLLLLCYFGLGKQTFKTVYGSWIFPVFI  
 KLTQSVPTLTHNSLLAALFGGVIVGCGLGIVFWSDSSTGGTGIIIQFLGKY  
 TPISLGQGVLIDGLVTIVGFLAFDSDTVMFSIIGLITISYIINAIQTGFTLST  
 VLIVSQEHQKIKTYINTVADRGVTEIPVKGGYSGTNQIMLMTTIAGYEFA  
 KLQEAIAEIDETAFITVTPTSQASGRGFSLQKNHGRLEDEDILMPM\*

ID-50

Clone 99

ATGAAAGAAAAACAGTCGAAAAGGCTTATTTATATACTACTGATTGTTCCCATTAT  
 CTTTATAAGTGTTTTTACATACAGTATTAGCCAGCCTTCTAAACTACTTCCACCAA  
 AGAATTAGTTATTCTAAGTCCAAATAGTCAAGCCATTTAACAGGAACGATTCCAG  
 CTTTTGAGGAAAAATACGGTATAAAAAGTTAAGCTTATTCAAGGTGGGACAGGGCA  
 ACTAATAGATAGATTAAGTAAGGAGGGTAAGCAGTTGAAGGCGGATATTTTCTTTG  
 GAGGAAATTATACGCAATTTGAAAGTCATAAGGCATTGTTTGAGTCTTACGTATCA

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FIG. 1 CONT'D

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AAGAATGTTCACTACTGTTATTCCAGACTATATCCATCCGAGTGATACGGCGACACC  
TTATACTATAAATGGGAGTGTCTTGATTGTAAATAACGAATTAGCTAAGGGACTTA  
CCATCAAGAGTTATGAAGATTTATTACAGCCTTCCTTAAAAGGTAAAATTGCCTTT  
GCAGATCCTCTAGAGTCGACCTGCAAGCATGCAAGCTTGGCGTAA

MKEKQSKRLIYILLIVPIIFISVFTYSISQPSKLLPPKELVILSPNSQAILTG TIPAFEKEYGI  
KVKLIQGGTGQLIDRLSKEGKQLKADIFFGGNYTQFESHKALFESYVSKNVHTVIPDYI  
HPSDTATPYTINGSVLIVNNELAKGLTIKSYEDLLQPSLKGKIAFADPLESTCKHASLA

ID-51

Clone 103

CCTCCTATCAAATGATGACAAACGTGAGAGGTACATGGAACAAATGCTCTTTAAAA  
TTGAAAATGCAACCTGGCAGCGTGTGGTAAGAGCACTTTATCGTAAATACAATAAG  
GAATTTTTTACATATCCAGCCGCCAAAACAAACCACCACGCTTTTGAATCAGGATT  
GGCATATCACACGGCAACAATGGTTCGTTTGGCAGATAGTATCGGAGATATCTATC  
CAGAACTTAATAAAAAGTTTGTATGTTTGGTGGTATTATGCTACATGATTTAGCCAAG  
GTCATAGAGTTATCGGGTCCTGATAATACAGAATATACTATTCGAGGTAATCTTAT  
CGGTCATATTTCACTTATTGATGAGGAATTA

LLSNDDKRERYMEQMLFKIENATWQRVVRALYRKNKEFFTYPAAKTNHHAFESGL  
AYHTATMVRLADSIGDIYPELNKSLMFAGIMLHDLAKVIELSGPDNTEY TIRGNLIGHIS  
LIDEEL

ID-52

Clone 104

ATGAAAAAAAAATAAAATTATCCGATTCAGTTTAGTTGGTGTCTACTT  
GCGATACTATGCTTTAGTCTTTTTGCTTTATTGAAGCCTAACAGTCAA  
CAATCATCATCTCAAAGTTGAGGAATGAGGATATAAAAAAGACATC  
CTCTCAAAAAAGAAATAAGAAATTACGATTACCAGCTGTATCATCAA  
AAGATTGGAACCTTGATTTTGGTCAATCGTGACCATAAACATGAAGAA  
TTAAGTCCAGATGTGGTGCCTGTTGAAAATATTTATTTGGATAAACGT  
ATTACGAAGCAAGCTACTCAGTTTTTAGAGGCTGCTAGAGCAATTGA  
TTCACGAGAACATTTAATTTTCGGGTATCGTAGTGTTGCCTATCAGGA  
GAAGTTGTTCAATTCTTATGTTACTCAAGAGATGACTAGTAACCCTAA  
TTTGACGAGGGGACAAGCAGAAAAGTTGGTAAAACTTACTCTCAGC  
CTGCAGGTGCTAGTGAACACCAGACTGGATTAGCGATGGATATGAGT  
ACTGTAGATTCTTTGAATGAGAGCGATCCTAGAGTAGTCAGTCAGTT  
GAAAAGATAGCTCCACAATATGGTTTTGTCTTACGGTTTCCGGATG  
GTAAAACAGCAGAAACAGGGGTAGGTTATGAAGATTGGCATTACCG

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FIG. 1 CONT'D

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CTATGTTGGGGTAGAGTCTGCAAAATATATGGTCAAACATCATTTAA  
CATTAGAAGAATACATAACTTTATTAAGGAGAATAACCAATGA

MKKNKIIRFSLVGVLLAILCFSLFALLKPNSQSSSQKLRNEDIKKTSSQK  
RNKKLRLPAVSSKDWNLILVNRDHKHEELSPDVVPVENIYLDKRITKQA  
TQFLEAARAIDSREHLISGYRSVAYQEKLFSYVTQEMTSNPNLTRGQA  
EKLVKTYSQPAGASEHQGLAMDMSVDSLNESDPRVVSQKKIAPQY  
GFVLRFPDGKTAETGVGYEDWHYRYVGVESAKYMKHHLTLEEYITLL  
KENNQ\*

ID- 53

Clone 106

CTGTTATGTGGATTTCTTCCATCAATTCCTGTGTCTAATTCCGGGGGG  
TATGGTATAATAACAGTTATGAAAAATAAAAAAATCTTATTTGGGAC  
TGGCCTTGCTGGTGTGGGTTTACTGGCAGCTGCTGGTTATACCCTAAC  
TAAAAAAGTAACAGATTATAAACGTCAGCAAATCACTCAGACCTTAA  
GAGAACTTTTTAGTCAGATGGGTGATATTCAGGTATTTTATTTAATG  
AATTTGAATCTGATATTAATAATGACCAGTGGTGGTCTTGTCTTGGAA  
GATGGCAGAATTTTCGAATTCATTTATCGTCAAGGTGTTCTTGATTAT  
GTGGAGGTGAGCAAATGA

LLCGFLPSIPVSNSSGGYGIITVMKNKKILFGTGLAGVGLLAAAGYTLTKK  
VTDYKRQQITQTLRELFQMGDIQVFYFNEFESDIKMTSGGLVLEDGRIF  
EFIYRQGVLDYVEVSK\*

ID-54

Clone 108

ATGTATCAAACCTCAGACAAATAAGGAAAAATTTGTTTTATTTTTGAAATTATTTATC  
CCAGTATTGATTTATCAATTTGCTAATTTTCAGCTACTTTTATTGATTCGGTTATGA  
CTGGACAGTATAGTCAGCTACATTTGGCAGGTGTGTCAACTGCTAGTAATTTATGG  
ACTCCGTTTTTTCGCTTTATTAGTAGGTATGATTTTCAGCATTAGTACCAGTAGTTGGT  
CAACATTTGGGTAGAGGAAATAAAGAACAAATTCGCACAGAATTTTCATCAATTTCT  
ATATTTAGGTTTGATACTGTCCTTAA

MYQTQTNKEKFVLFKLFIPVLIYQFANFSATFIDSVMTGQYSQLHLAGVSTASNLWTP  
FFALLVGMISALVPVVGQHLGRGNKEQIRTEFHQFLYLGLLISL

ID-55

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FIG. 1 CONT'D



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 Clone 112

CTGCTCTTTTTAGCTAACTTTTCTAATTTATGGTATAATTGTATGGATT  
 GTTTAGCTAGAATGGAGAAGATGATGCAAGATGTTTTTCATTATAGGA  
 AGTAGAGGGTTGCCAGCTCGTTACGGTGGTTTTGAACTTTTGTTC  
 GAATTGATTAATCATCAAAAAAGTTCCGACATAAAATACCATGTTGC  
 ATGCCTTAGTGATAAAGAACATCATACTCATTTTAACTTTGCTGACGC  
 TGATTGTTTTACTATAAATCCTCCCAATTAGGGCCAGCACGTGTGAT  
 TGCTTATGATATTATGGCCATTAATTATGCCCTTGACTTGGTTAAGAC  
 ACATGATTTAAAAGAGCCTATTTTTTATATTTTAGGAAATACAATTGG  
 TGCCTTTATTTGGCATTGTTGCCAATAAAATACATAAAGTCGGTGGCTT  
 ATTGTATGTTAATCCGGATGGTTTAGAGTGAAGCGATCAAAGTGGT  
 CTCGTCCCACACAGCGTTATTTAAAATACGCCGAAAAATGTATGACT  
 AAAAATGCAGACCTAATTATTTCTGATAATATTGGTATTGAAAATTA  
 CATTCAATCTACCTACTCTAATGTGAAGACAAGGTTTCATTGCTTACGG  
 TACAGAGATTAATTCTAGGAAATTATCGTCAGATGATCCACGTGTCA  
 AACAGTTGTTTAAAAAATGGAATATTAAGTCTAAGGGTACTATCTA  
 ATCGTTGGTCGATTTGTCCCTGAAAACAATTATGAAACGGCTATTAG  
 GGAGTTCATGGCTTCAGATACTAAGCGTGATTTAGTTATTATCTGTAA  
 CCATCAAAATAAACCCTACTTTGAAAAGTTGTCCTTAAAGACAAACC  
 TTCAACAAGATAAAAGAGTTAAGTTTGTAGGTACGCTCTATGAAAAA  
 GATCTGCTGGATTATGTTTCGTCAACAAGCCTTTGCTTATATTCATGGG  
 CATGAAGTTGGCGGTACTAATCCAGGACTGCTTGAGGCTTTAGCTAA  
 TACTGATTTGAATCTTGTTCTAGATGTTGATTTCAACAAATCAGTAGC  
 AGGTCTCTCAAGTTTTTACTGGACTAAAAAAGAGGGGGATTTAGCTA  
 AGCTT

MLFLANFSNLWYNCMDCLARMEKMMQDVFIIGSRGLPARYGGFETFVS  
 ELINHQKSSDIKYHVACLSDKEHHTHFNFADADCFTINPPQLGPARVIAY  
 DIMAINYALDLVKTHDLKEPIFYILGNTIGAFIWHFANKIHKVGGLLYVN  
 PDGLEWKRKWSRPTQRYLKYAEKCMTKNADLIISDNIGIENYIQSTYSN  
 VKTRFIA YGTEINSRKLSSDDPRVKQLFKKWNKSKGYLIVGRFVPENN  
 YETAIREFMASDTKRDLVIICNHQNNPYFEKLSLKTNLQQDKRVKFGVT  
 LYEKDLLDYVRQAFAYIHGHEVGGTNPGLLEALANTDLNLVLDVDFN  
 KSVAGLSSFYWTKKEGDLAKL

ID-56

## Clone 120

TTGAGGAGTAATATGGTAAAGACAGCAGTTTTAATGGCGACATACAA  
 TGGCGAAAAATTTATATCTGAACAACCTTGATTCAATTCGCCAACAGA  
 CATAAAACCAGATTATGTATTATTGAGGGATGATTGTTCAACGGAT  
 GAAACAGTCAATGTCGTCAATAACTATATCGCAAAACATGAGTTAGA

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FIG. 1 CONT'D

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AGGCTGGAAAATTGTAAAAACGACAAAACTTAGGCTGGCGTTTAA  
ATTTTCGTCAATTACTTATTGATGTGTTAGCCTATGAGGTTGACTATG  
TCTTTTTTAGTGATCAAGATGATATTTGGTATCTTGATAAAAAACGAAC  
GACAGTTTGCCATTATGTCAGATAACCCTCAAATTGAGGTTTTGAGTG  
CAGACGTTGATATCAAACGATGTCTACAGAAGCCAGTGTTCCACAT  
TTTCTAACTTTTTCTTCTAGTGATAGAATCAGTCAGTATCCTAAAGTA  
TATGATTATCAAACATTCCGTCCCGGATGGACCATTGCTATGAAGAG  
AGATTTTGCGCAAGCTATCGCTTGA

MRSNMVKTAVLMATYNGEKFISEQLDSIRQOTLKP DYVLLRDDCSTDET  
VNVVNNYIAKHELEGWKIVKNDKNL GWRLNFRQLLIDVLAYEVDYVFF  
SDQDDIWYLDKNERQFAIMSDNPQIEVLSADVDIKTMSTEASVPHFLTFS  
SSDRISQYPKVYDYQTFRPGWTIAMKRDFQAIA\*

ID-57

Clone 123

GTGATTATGGATAAGTCTATTCCTAAAGCAACTGCTAAACGTTTATCA  
CTGTACTACCGTATTTTTAAACGTTTTAATACTGATGGCATCGAAAAA  
GCTAGTTCCAAACAAATTGCAGATGCCCTAGGTATCGATTCTGCTACT  
GTTTCGACGTGATTTTTCTTATTTTGGTGAAGTAGGACGCCGTGGTTTT  
GGTTATGATGTCAAAAACTTATGAACTTCTTTCAGAAATATTGAA  
CGATCATTCTACAACAAATGTTATGCTGGTGGGGTGTGGAAATATCG  
GTAGAGCTCTCTTGCATTATCGTTTCCACGATCGCAATAAAATGCAA  
ATTTCAATGGCTTTTGATTTAGATAGCAATGATTTAGTTGGTAAAACA  
ACCGAGGATGGAATTCCTGTCTACGGTATTTTCGACTATCAATGACCA  
TTAATAGATAGTGATATTGAACTGCTATCCTAACAGTACCTAGTAC  
AGAAGCCCAAGAAGTTGCTGACATCTTAGTCAAAGCAGGTATAAAA  
GGCATCTTGAGTTTTTCTCCAGTTCATTTAACATTACCAAAGATATC  
ATTGTTCAAGTATGTAGATTTAACAAGCGAATTACAACTTTACTTTAT  
TTCATGAACCAGCAGCGATAA

MIMDKSIPKATAKRLSLYRIFKRFNTDGIEKASSKQIADALGIDSATVRR  
DFS YFGELGRRGFGYDVKKLMNFFAEILNDHSTTNVMLVGCNIGRALL  
HYRFHDRNKMQISMAFDLDSNDLVGKTTEDGIPVYGISTINDHLIDSDIE  
TAILTVPSTEAQEVADILVKAGIKGILSFVHLTLPKDIIVQYVDLTSELQ  
TLLYFMNQQR\*

ID-58

Clone 125

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FIG. 1 CONT'D

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ATGGGTGCTAAAGGAGCAGATGTCATTCTCGTTTTATCACACTCTGGCATTGGAGA  
TGATCGATATGAAGAAGGTGAAGAAAACGTTGGCTATCAAATTGCCAGCATCAAG  
GGAGTGGATGCCGTTGTTACGGGACACTCACACGCTGAATTTCCATCAGGTAACGG  
TACTGGCTTCTATGAAAAATACACTGGAGTTGATGGTATCAATGGAAAAATAAATG  
GAACACCTGTTACAATGGCAGGCAAGTACGGGGATCACCTTGGTATTATTGATTTA  
GGACTTAGTTATACTAATGGAAAATGGCAAGTCTCCGAAAGCAGTGCTAAAATCC  
GTAAAATTGATATGAACTCAACAACCTGCTGACGAGCGTATCATTGCATTGGCTAAG  
GAAGCACACGATGGCACTATCAACTATGTTTCGCCAACAAGTAGGTACAACAACCTG  
CGCCAATTACAAGTTACTTTGCACTAGTTAA

MGAKGADVILVLSHSGIGDDRYEEGEENVGYQIASIKGVDAVVTGHSHAEPSPGNGTG  
FYEKYTGVDGINGKINGTPVTMAGKYGDHLGIIDLGLSYTNGKWQVSESSAKIRKIDM  
NSTTADERIILAKEAHDGTINYVRQQVGTTPITSYFALV

ID-59

Clone 135

TTGTCAATAAGGTTTCAAATCAGCTTGAAATATGATAAAAATAAAACAGATTGTAAG  
TGACTGTTTAAGCTTGTTTTTTCAGAGAGGTTTTTATGAATACAAACACAATAAAAA  
AGGTTGTAGCGACTGGAATTGGAGCTGCACTTTTTATCATTATAGGTATGCTAGTT  
AA

MSIRFQISLKYDKIKQIVSDCLSLFFREVMNTNTIKKVVATGIGAALFIIIGMLV

ID-60

Clone 145

ATGAAACATTTAAAATTTCAATCGGTCTTCGACATTATTGGTCCTGTTATGATTGGA  
CCATCAAGTAGTCATACTGCAGGAGCTGTCCGCATTGGTAAAGTTGTCCATTCTAT  
TTTTGGTGAACCTAGTGAAGTAACCTTTCATTTATACAATTCTTTTGCTAAAACCTA  
CCAAGGACACGGTACTGATAAAGCATTGGTTGCAGGGATTCTAGGAATGGATACA  
GATAATCCAGATATTAA

MKHLKFQSVFDIIGPVMIGPSSSHTAGAVRIGKVVHSIFGEPSEVTFHLYNSFAKTYQG  
HGTDKALVAGILGMDTDNPD

ID-61

Clone 147

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FIG. 1 CONT'D

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GTGTCAGAAGGTGTTTTAATGTTTCTAAAAGAAGATGACGTAGAGACTTTTCTTCA  
TATCCTGACAAATTCATTTAGCCAATTTATGGCACAATTTGATTTGTGTCATAAGGA  
AATGATTAA

ID-62

Clone 150

ATGACCTACAAAGATTACACAGGTTTAGATCGGACTGAACTTTTGAGTAAAGTGCG  
TCATATGATGTCCGACAAACGTTTTAA

MTYKDYTGDLRTELLSKVRHMMSDKRF

ID-63

Clone S2

CTGAGTTGGGTCTTGGAACGGTCCTGTCAATCATACTAGCTATCAAGGAGACTAA  
AATGTATTTAGAACAATAAAAGAGGTAAATCCTTTAA

MSWVLETVLSIILAIKETKMYLEQLKEVNPL

ID-67

Clone 3-40

GTGAAAAAAAAAATTAGTCTCATCACTTCTAAAGTGTTCTCTAATCATT  
ATTGTTAGCTTTGCTGGTGGAGCATTGCTAGTTTTGTCATGAATCAT  
AATGACAATATTCCAAATGGTGGTGTCACTAAAAGTAAAGTAAA  
TTATAATAACATAACGCCTACAACAAAAGCTGTAAAAAGGTACAAA  
ATAGTGTTGTTTCTGTTATCAATTATAACAACAAGAGAGTCGTTCTG  
ACCTATCAGACTTCTATAGTCATTTTTTTGGTAATCAGGGGGGCAACA  
CTGATAAGGGCTTACAAGTTTACGGTGAAGGCTCTGGAGTCATCTAT  
AAAAAAGATGGTAAAAATGCCTATGTTGTCACTAATAACCACGTCAT  
TGATGGGGCTAAACAAATTGAAATTCAACTAGCTGATGGCTCAAAAG  
CAGTTGGGAAACTTGTTGGGTCAGATACCTACTCTGATTTAGCCGTCG  
TCAAATTCATCAGATAAAGTTTCAAATATTGCAGAATTTGCTGATT  
CATCAAAACTCAACATTGGTGAAACTGCTATAGCGATCGGAAGCCCT  
CTTGGAAGTGAATGCAAATTCTGTAAGTCAAGGTATTGTATCTAGT  
TTAAAAAGAACTGTAACAATGACTAATGAAGAAGGACAAACAGTTT  
CTACAAATGCTATCCAGACGGATGCTGCTATCAATCCTGGTAATTCA  
GGTGGAGCACTTATCAATATTGAAGGACAGGTTATTGGAATTAATTC  
TAGTAAAATTTCTTCTACATCAAATCAAACCTCAGGACAATCGTCAG

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FIG. 1 CONT'D

GAAATAGCGTTGAAGGTATGGGATTTGCCATTCCTTCAAATGATGTT  
 GTTAAGATTATCAATCAACTTGAGAGTAACGGACAAGTAGAGAGACC  
 TGCTCTAGGTATTTCTATGGCTGGATTAAGTAATTTACCATCCGATGT  
 TATTAGTAAACTGAAAATCCCAAGTAATGTTACTAATGGTATTGTAG  
 TAGCATCTATCCAATCTGGCATGCCAGCTCAAGGCAAACCTAAAGAAA  
 TACGATGTCATTACTAAAGTTGACGATAAAGAAGTAGCATCTCCAAG  
 TGATTTACAAAGTTTACTCTATGGCCACCAGGTAGGGGATTCCATAA  
 CAGTAACCTTTTATCGTGGTGAAAATAAACAAACAGTCACTATAAAA  
 CTTACTAAACTAGTAAAGATTTAGCTAAACAACGAGCAAATAACTA  
 A

MKKKLVSSLLKCSLIIIVSFAGGAFASFVMNHNDNIPNGGVTKTSKVNY  
 NNITPTTKAVKKVQNSVVSVINYKQESRSDLDFYSHFFGNQGGNTDK  
 GLQVYGEESGVIYKKDGKNA YVVTNNHVIDGAKQIEIQLADGSKAVGK  
 LVGSDTYSDLA VVKIPSDKVSNIAEFADSSKLNIGETAIAIGSPLGTEYAN  
 SVTQGIVSSLKRTVTMTNEEGQTVSTNAIQTDAAINPGNSGGALINIEGQ  
 VIGINSSKISSTSNQTSQSSGNSVEGMGFAIPSNDVVKIINQLESNGQVE  
 RPALGISMAGLSNLPDVISKLIKPSNVTNGIVVASIQSGMPAQGKLLKY  
 DVITKVDDEKVASPSDLQSLLYGHQVGDSTVTFYRGENKQVTIKLTKT  
 SKDLAKQRANN\*

ID-68

Clone 3-30

ATGTTAAAATGGTATACAAACAAAGGAGGGAGGATGATAATGAAGA  
 AATGTTTTTTGGCTATTTGTTTAGCTCTTAGTTTTTTTATGGTTTCAGT  
 TCAAGCAGATGAGGTGGACTATAACATTCCTCATTATGAGGGTAATC  
 TAACTATTCACAATGATAATAGTGCTGATTTTACAGAGAAGGTTACTT  
 ACCAATTTGATTCGTCCTATAATGGACAGTATGTCACGTTAGGTACG  
 GCGGGTAAGTTATCTGACAATTTTGATATTAATAATAAGCCACAGGT  
 TGAAGTTTCAATTAATGGTAAAGTAAGGAAAGTTAGTTACCAGATAG  
 AAGATTTGGAGGATGGCTACCGTTTGAAAGTGTTAATGGTGGTGAA  
 GCAGGTGATACTGTTAAAGTCAATGTTTCAGTGGAAACTAAAAAATGT  
 TCTATTTATGCATAAGGATGTTGGTGAACCTAACTGGATTCCTATTAG  
 CGACTGGGATAAAACGTTAGAGAAAGTAGATTTTTGGATATCAACTG  
 AAAAAAGGTTGCTCTTTCTCGTCTTTGGGGGCACTTGGGTTATCTTA  
 AACTCCTCCTAAAATAAGACAAAATAATAATCGTTACCATTTGACA  
 GCTTTAATGTAAACAAACGATTAGAATTTTCATGGTTATTGGGATAG  
 ATCTTATTTAATCTACCTACAAACAGTAAAAATAATTACAAGAAAA  
 AAATTGAACATCAAGAGAAGATAATAGAGCGTCATGGTTTTATCCTA  
 AGTTTCTTGTTAAGGATATTATTACCTTCATTCTTTATTATTGTGACAC  
 TATTCATCTCAATTAGGGTGTTTCCTGTTTAGAAAAAAGTTAATAAAT

FIG. 1 CONT'D

ACGGGCAATTCCTAAGGATCATCATTATATGAAGCACCTGAGGAC  
 CTTTCACCATTAGAGTTAACTCAAAGCATTATAGTATGAGCTTTAAA  
 AATTTTCAAGATGAGGAGAAGAAAACCTCACCTTATCAGTCAAGAACA  
 ACTCATAACAGTCAATTCTATTAGACTTGATTGATAGAAAAGTATTGA  
 ATTATGATGATAACTTGTTATCTCTAGCTAACTTAGATAGAGCTTCTG  
 ATGCAGAAATAGATTTTATAGAGTTTGCTTTTGCGGATTCTACGAGTT  
 TGAAGCCAGATCAACTCTTTTCTAATTACCAATTTAGTTATAAAGAAA  
 CACTACGTGAACTGAAAAAGCAGCACAAGGCTTCAGATCTGCAAAAT  
 CAAATGAGACGCCGAGGAAGTAATGCCTTATCAAGAATTACGCGTCT  
 CACAAGGTTGATTTCTAAAGACAATATAAACTCTCTTAGAAGAAAGG  
 GAATTCATCCCCTTATCGTAAAATGTCTTCAGAAGAGTCTAAAGAA  
 TTATCTAGGTTAAAAAGATTACGTTACCTATCACCTCTTATTTCTTTTG  
 TTGTTATAATTTATACGCTTTTTTTAAATTATTTACCTATTTCTGTAT  
 CTATCTCTTATTGTTTGGTGTATCCTGTTGTTGAATAAAAATCATT  
 ATGATGACAAGAAAAATAAGTAACGGTTATATTGTAAGTGAAGATGG  
 AGCAAGTCGTGTCTACCAATGGACTAGTTTTAGGAACATGCTAAGGG  
 ATATCAAATCGTTTGATCGTTCAGAGTTAGAAAGTATCGTATTATGG  
 AATCGAATATTGGTTTACGCTACTTTATTCGGCTACGCTGACCGTGT  
 GAGAAAGTACTCAGAGTGAACCAATAGATATTCCAGAAAGATTTGC  
 AACATTGATAGTCATCGATTTGCGATTTACAGTCAATCAATCTAGTAA  
 TCATTTTTCAACGATAACTGAAGATGTTAGTCACGCTTCTAATTTTAG  
 TGTTAATTCAGGCGGTTCTTCAGGTGGTTTCTCAGGCGGCGGAGGCG  
 GCGGAGGTGGCGGTGCCTTCTAA

MLKWYTNKGGRMIMKKCFLAICLALSFFMVSVQADEVDYNIPHYEGLN  
 TIHNDNSADFTEKVYQFDSSYNGQYVTLGTAGKLSDNFDINNKPQVEV  
 SINGKVRKVSQIEDLEDGYRLKVFNGGEAGDTVKNVQWKLKNVLF  
 MHKDVGELNWPISDWDKTLKVFDFWISTDKKVALSRLWGHLGYLKTP  
 PKIRQNNRYHLTAFNVNKRLEFHGYWDRSYFNLPTNSKNNYKKKIEH  
 QEKIHERHGFI LSFLLRILLPSFFIIVTLFISIRVFLFRKKVNKYGQFPKDHHL  
 YEAPEDLSPELQTQSIYSMSFKNFQDEEKKTHLISQEQLIQSILLDLDRKV  
 LNYDDNLLSLANLDRASDAEIDFIEFAFADSTSLKPDQLFSNYQFSYKET  
 LRELKKQHKASDLQNQMRRRGSNALSRLRTRLSKDNINSLRRKGISS  
 PYRKMSSEESKELSRKRFSYLSPLISFVVIYTLFLNYFTYFCIYLLLFVVI  
 LLLNKIIFMMTRKISNGYIVTEDGASRVYQWTSFRNMLRDIKSFDRESELE  
 SIVLWNRILVYATLFGYADRVEKVLVRVNQIDIPERFANIDSHRFAISVNQS  
 SNHFSTITEDVSHASNFSVNSGGSSGGFSGGGGGGGGAF\*

ID-69

Clone 3-38

ATGATGATTGTGAATAATGGTTATCTAGAAGGGAGAAAAATGAAAA  
 AGAGACAAAAAATATGGAGAGGGTTATCAGTTACTTTACTAATCCTG

FIG. 1 CONT'D

TCCCAAATTCCATTTGGTATATTGGTACAAGGTGAAACCCAAGATAC  
CAATCAAGCACTTGGAAAAGTAATTGTTAAAAAACGGGAGACAAT  
GCTACACCATTAGGCAAAGCGACTTTTGTGTTAAAAAATGACAATGA  
TAAGTCAGAAACAAGTCACGAAACGGTAGAGGGTTCTGGAGAAGCA  
ACCTTTGAAAACATAAAACCTGGGAGACTACACATTAAGAGAAGAAA  
CAGCACCAATTGGTTATAAAAAAACTGATAAAACCTGGAAAGTTAAA  
GTTGCAGATAACGGAGCAACAATAATCGAGGGTATGGATGCAGATA  
AAGCAGAGAAACGAAAAGAAGTTTTGAATGCCCAATATCCAAAATC  
AGCTATTTATGAGGATACAAAAGAAAATTACCCATTAGTTAATGTAG  
AGGGTTCCAAAGTTGGTGAACAATACAAAGCATTGAATCCAATAAAT  
GAAAAGATGGTTCGAAGAGAGATTGCTGAAGGTTGGTTATCAAAAA  
AAAATCCAGGGGTCAATGATCTCGATAAGAATAAATAAAAATTGAA  
TTAACTGTTGAGGGTAAAACCACTGTTGAAACGAAAGAAGTTAATCA  
ACCACTAGATGTCGTTGTGCTATTAGATAATTCAAATAGTATGAATA  
ATGAAAGAGCCAATAATTCTCAAAGAGCATTAAAAGCTGGGGAAGC  
AGTTGAAAAGCTGATTGATAAAATTACATCAAATAAAGACAATAGA  
GTAGCTCTTGTGACATATGCCTCAACCATTTTGTATGGTACTGAAGCG  
ACCGTATCAAAGGGAGTTGCCGATCAAATGGTAAAGCGCTGAATG  
ATAGTGTATCATGGGATTATCATAAACTACTTTTACAGCAACTACA  
CATAATTACAGTTATTTAAATTTAACAAATGATGCTAACGAAGTTAA  
TATTCTAAAGTCAAGAATTCCAAAGGAAGCGGAGCATATAAATGGG  
GATCGCACGCTCTATCAATTTGGTGCACATTTACTCAAAAAGCTCTA  
ATGAAAGCAAATGAAATTTTAGAGACACAAAGTTCTAATGCTAGAAA  
AAAACCTATTTTTCACGTAAGTGGTGTCCCTACGATGTCTTATGC  
CATAAATTTTAATCCTTATATATCAACATCTTACCAAACCAAGTTAA  
TTCTTTTTTAAATAAAAATACCAGATAGAAGTGGTATTCTCCAAGAGG  
ATTTTATAATCAATGGTGATGATTATCAAATAGTAAAAGGAGATGGA  
GAGAGTTTTAACTGTTTTCGGATAGAAAAGTTCCTGTTACTGGAGG  
AACGACACAAGCAGCTTATCGAGTACCGCAAATCAACTCTCTGTAA  
TGAGTAATGAGGGATATGCAATTAATAGTGGATATATTTATCTCTATT  
GGAGAGATTACAAGTGGGTCTATCCATTTGATCCTAAGACAAAGAAA  
GTTTCTGCAACGAAACAAATCAAACCTCATGGTGAGCCAACAACATT  
ATACTTTAATGGAAATATAAGACCTAAAGGTTATGACATTTTTACTGT  
TGGGATTGGTGTAACGGGAGATCCTGGTGCAACTCCTCTTGAAGCTG  
AGAAATTTATGCAATCAATATCAAGTAAAACAGAAAATTATACTAAT  
GTTGATGATACAAATAAAATTTATGATGAGCTAAATAAAACTTTAA  
ACAATTTGTTGAGGAAAAACATTCTATTGTTGATGGAAATGTGACTG  
ATCCTATGGGAGAGATGATTGAATTCCAATTAATAAATGGTCAAAGT  
TTTACACATGATGATTACGTTTTGGTTGGAAATGATGGCAGTCAATTA  
AAAAATGGTGTGGCTCTTGGTGGACCAAACAGTGATGGGGGAATTTT  
AAAAGATGTTACAGTGACTTATGATAAGACATCTCAAACCATCAAAA  
TCAATCATTGAACTTAGGAAGTGGACAAAAAGTAGTTCTTACCTAT  
GATGTACGTTTTAAAGATAACTATATAAGTAACAAATTTTACAATAC  
AATAATCGTACAACGCTAAGTCCGAAGAGTGAAAAAGAACCAAAT

FIG. 1 CONT'D

ACTATTCGTGATTTCCCAATTCCCAAATTCGTGATGTTTCGTGAGTTT  
 CCGGTACTAACCATCAGTAATCAGAAGAAAATGGGTGAGGTTGAATT  
 TATTAAGTTAATAAAGACAAACATTCAGAATCGCTTTTGGGAGCTA  
 AGTTTCAACTTCAGATAGAAAAAGATTTTTCTGGGTATAAGCAATTT  
 GTTCCAGAGGGAAGTGATGTTACAACAAAGAATGATGGTAAAATTTA  
 TTTTAAAGCACTTCAAGATGGTAACTATAAATTATATGAAATTTCAA  
 GTCCAGATGGCTATATAGAGGTTAAAACGAAACCTGTTGTGACATTT  
 ACAATTCAAAAATGGAGAAGTTACGAACCTGAAAGCAGATCCAAATG  
 CTAATAAAAATCAAATCGGGTATCTTGAAGGAAATGGTAAACATCTT  
 ATTACCAACTCCCAAACGCCACCAGGTGTTTTTCTAAAACAGG  
 GGAATTGGTACAATTGTCTATATATTAGTTGGTTCTACTTTTATGAT  
 ACTTACCATTTGTTCTTTCCGTCGTAAACAATTGTAA

MMIVNNGYLEGRKMKKRQKIWRGLSVTLLILSQIPFGILVQGETQDTNQ  
 ALGKVVVKKTGDNATPLGKATFVLKNDNDKSETSHETVEGSGEATFENI  
 KPGDYTLREETAPIGYKKTDKTWKVKVADNGATIIEGMDADKAEKRKE  
 VLNAQYPKSAIYEDTKENYPLVNVEGSKVGEQYKALNPINGKDGREIA  
 EGWLSKKNPGVNDLDDKNKYKIELTVEGKTTVETKELNQPLDVVLLDN  
 SNSMNNERANNSQRALKAGEAVEKLIDKITSNKDNRVALVTYASTIFDG  
 TEATVSKGVADQNGKALNDSVSWDYHKTTFTATTHNYSYLNLTNDAN  
 EVNILKSRIPEAEHINGDRTLQYFGATFTQKALMKANEILETQSSNARK  
 KLIFHVTDGVPTMSYAINFNPISTSYQNQFNSFLNKIPDRSGILQEDFIIN  
 GDDYQIVKGDGESFKLFSRDKVPVTGGTTQAAAYRVPQNQLSVMSNEGY  
 AINSGYIYLWWRDYNWVYFPDPKTKKVSATKQIKTHGEPTTLYFNGNIR  
 PKGYDIFTVGIGVNGDPGATPLEAEKFMQSISSTENYTNVDDTNKIYDE  
 LNKYFKTIVEEKHSIVDGNVTDPMGEMIEFQLKNGQSFTHDDYVLVGNL  
 GSQKNGVALGGPNSDGGILKDVTVTYDKTSQTIKINHLNLGSGQKVVL  
 TYDVRLKDNYSNKFYNTNRTTTLSPKSEKEPNTIRDFPIPKIRDVREFPV  
 LTISNQKMGVEFEIKVNKDKHSESLGAKFQLQIEKDFSGYKQFVPEGS  
 DVTTKNDGKIYFKALQDGNKYLYEISSPDGYIEVKTKPVVTFITQNGEVT  
 NLKADPNANKNQIGYLEGNGKHLITNTPKRPPGVFPKTGGIGTIVYILVG  
 STFMILTICSFRRKQL\*

ID-70

Clone 141

ATGAATAGAAAAGTTGAGGAAAAAATGGCTGGGAATCGTAATAACG  
 ATATGAATGTCTATTGTTCAATTTGTGGCAAAGCCAAGATGAAGTA  
 AAAAAAATTATTGCAGGTAATGGTGTTCATTTGTAATGAATGTGTG  
 GCCTTATCACAAGAAATTATTAAGGAAGATTAGCTGAGGAAGTACT

FIG. 1 CONT'D



GGCTCATTTAGCAGAAGTACCAAAACCTAAGGAACTATTAGAAATAT  
 TAAATCAATATGTTGTAGGGCAAGATCGTGCTAAACGTGCTTTAGCA  
 GTTGCTGTCTACAATCATTACAAGCGTGTTAGTTATACCGAGAGTAGT  
 GACGATGATGTAGATTTGCAAAAATCCAACATTTTGATGATTGGTCC  
 AACTGGCTCAGGAAAACCTTCTTAGCACAACACTGGCTAAAAGCC  
 TTAATGTACCGTTTGCTATTGCAGATGCGACTTCATTGACCGAAGCAG  
 GATACGTTGGAGAAGATGTTGAGAATATTCTTCTTAAATTGATTCAA  
 GCTGCTGATTATAATGTGCAACGTGCTGAGCGTGGTATTATCTACGTT  
 GATGAAATAGATAAAATTGCTAAGAAAGGCGAAAATGTTTCTATCAC  
 ACGTGATGTGTCTGGTGAAGGTGTACAGCAAGCCCTTCTTAAAATTA  
 TTGAGGGTACGGTAGCAAGTGTTCCCCCACAGGGTGGGCGTAAACAT  
 CCTAACCAAGAAATGATTCAAATTAATACCAAGAACATCCTTTTTTATT  
 GTCGGTGGTGCTTTTGATGGTATTGAAGACCTTGTGAAGCAACGTTTA  
 GGCGAAAAGTTATTGGTTTTGGACAGACAAGCCGTAAAATTGATGA  
 CAACGCTTCTTATATGCAAGAGATAATTTCTGAGGATATTCAAAGT  
 TTGGACTGATTCCAGAGTTTATTGGCCGTTTACCAGTAGTTGCAGCGT  
 TAGAACTTCTTACTGCAGAAGATCTGGTTCGTATTCTGACAGAACCA  
 CGCAATGCTTTGGTTAAACAATACCAAACCTTATTATCTTATGATGGT  
 GTAGAATTGGAATTTGACCAGGATGCTCTATTGGCTATCGCTGATAA  
 GGCTATCGAGCGCAAGACTGGTGCACGTGGTTTTACGTTCTATTATTG  
 AAGAAACGATGCTTGATATCATGTTTGAAATTCCAAGCCAAGAAGAT  
 GTAACAAAAGTTCGTATCACAAAGGCTGCTGTTGAGGGTACTGACAA  
 GCCTGTTTTAGAGACGGCTTAG

MNRKVEEKMAGNRNNDMNVYCSFCGKSQDEVKIIAGNGVFICNECV  
 ALSQEIKEELAEVLAHLAEVPKPKELEILNQYVVGQDRAKRALAVA  
 VYNHYKRVSYTESSDDDDVDLQKSNILMIGPTGSGKTFLAQTAKSLNVP  
 FAIADATSLTEAGYVGEDVENILKLIQAADYNVERAERGIYVDEIDKIA  
 KKGENV SITRDVSGEGVQQALLKIIEGTVASVPPQGRKHPNQEMIQINT  
 KNILFIVGGAFDGIEDLVKQRLGEK VIGFGQTSRKIDDNASYMQEIISEDI  
 QKFGLIPEFIGRLPVVALELLTAEDLVRILTEPRNALVKQYQTLLSYDG  
 VELEFDQDALLAIADKAIERKTGARGLRSIIETMLDIMFEIPSQEDVTKV  
 RITKAAVEGTDKPVLETA\*

ID-71

Clone 3-20

ATGAAAAGATTACATAAACTGTTTATAACCGTAATTGCTACATTAGG  
 TATGTTGGGGGTAATGACCTTTGGTCTTCCAACGCAGCCGCAAACG  
 TAACGCCGATAGTACATGCTGATGTCAATTCATCTGTTGATACGAGC  
 CAGGAATTTCAAATAATTTAAAAAATGCTATTGGTAACCTACCATT  
 TCAATATGTTAATGGTATTTATGAATTAATAATAATCAGACAAATTT  
 AAATGCTGATGTCAATGTTAAAGCGTATGTTCAAATAACAATTGACA

FIG. 1 CONT'D

---

ATCAACAAAGACTATCAACTGCTAATGCAATGCTTGATAGAACCATT  
 CGTCAATATCAAAATCGCAGAGATACCACTCTTCCCGATGCAAATTG  
 GAAACCATTAGGTTGGCATCAAGTAGCTACTAATGACCATTATGGGC  
 ATGCAGTCGACAAGGGGCATTTAATTGCCTATGCTTTAGCTGGAAAT  
 TTCAAAGGTTGGGATGCTTCCGTGTCAAATCCTCAAAATGTTGTCACA  
 CAAACAGCTCATTCCAACCAATCAAATCAAAAATCAATCGTGGACA  
 AAATTATTATGAAAGCTTAGTTCGTAAGGCGGTTGACCAAAACAAAC  
 GTGTTTCGTTACCGTGTAACTCCATTGTACCGTAATGATACTGATTTAG  
 TTCCATTTGCAATGCACCTAGAAGCTAAATCACAAGATGGCACATTA  
 GAATTTAATGTTGCTATTCCAACACACAAGCATCATACTACTATGGA  
 TTATGCAACAGGAGAAATAACACTAAATTAA

MKRLHKLFIATLGM LGVMTFGLPTQPQNVTPIVHADVNSSVDTSQE  
 FQNNLKNAIGNLFPQYVNGIYELNNNQTNLNADVNVKAYVQNTIDNQQ  
 RLSTANAMLDRTIRQYQNRDRTLDPANWKPLGWHQVATNDHYGHAV  
 DKGHLIAYALAGNFKGWDASVSNPQNVVTQTAHSNQSNOQKINRGQNY  
 YESLVRKAVDQNKRVRYRVTPLYRNDTDLVPFAMHLEAKSQDGTLEFN  
 VAIPNTQASYTMDYATGEITLN\*

ID-72

Clone 13

ATGAAAACTATCGAAAACTTATTGTACTACTACTTCTAATCTTTTTT  
 GCCATTTTTATGGGAGCATATGCTTACACGCATATTGTTGAAAAAAG  
 ATCCCTAACTAGCAATACTATTGAAAAACTCTACCTGTGGTAAATC  
 AGATTAAGCCTCAAACCATTAAAGAATACCAAAATTACTTAATAAG  
 GTAGCTAAACGTAATGTTCTTCTGTAGACATTCCTCAGGCATTAAT  
 AATGAAAAGGTAGAAATTACTGCTACTGATGGCATGCAAACATTAC  
 TTGGAATGATAAAAATAATCCTAAGCAAAAGGTTATCTTCTATGTTT  
 ATGGAGGATCATATATCCATCAAGCTTCCGAATTACAATATATTTTTG  
 TCAATAAACTAGCTAAAAAATTAGATGCAAAAGTTGTCTTTCCTATTT  
 ACCCTAAAGCTCCTACATATAATTATAGTGATGCTATCCCCAAAATTA  
 AAAAATTATACCAAAATACATTAGCTAGCGTCACATCTCACAAACAG  
 ATTATCCTAGTAGGTGAAAGTGCAGGCGGAGGCCTTGCTTTAGGTAT  
 TGCTGATAACCTTGCACGGAGCATATCAAACAACCAAAAGAAATTAT  
 TTAA

MKNYRKLIVLLLLIFFAIFMGAYAYTHIVEKRSLTSNTIEKTLPVVNQIKP  
 QTIKEYQNYLTKVAKRNVLPVDIPQALNNEKVEITATDGMQFTWNDK  
 NNPKQKVIFVYVHGGSYIHQASELQYIFVNKLAKKLDKVVFPPIYPKAPT  
 YNYSDAIPKIKKLYQNTLASVTSHKQIILVGESAGGGLALGIADNLARSIS  
 NNQKKLF\*

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FIG. 1 CONT'D

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 ID-73

Clone 2-19

TTGATTCTAATAACTTCCTATGGGATAATATCTTTATCACAAAAATTG  
 AGGGAATTTATTATGAAGTTAAAACATATTGTCTTAGGATTAGCCTTA  
 ACAACACTTTTAGGAGTCACATTTAGTAATCAAGAAGTTTCAGCAAG  
 CTCAACTTCAAGTAAAGTTGTTAAAGTTGGTGTATGACCTTTTCTGA  
 CACTGAAAAAGCACGTTGGGATAAAATTGAAAAGCTAGTAGGTGAT  
 AAAGCTAAAATCAAATTTACAGAATTTACAGATTATACACAACCAAA  
 TCAAGCGACAGCCAATAAGGATGTGGATATTAATGCCTTTCAACATT  
 ACAATTTCTTAGAAAAGCTGGAATAAGGAAAATAAGAAAAACTTAATT  
 CCACTTGAAAAGACTTACTTAGCTCCAATTCGTATCTATTCTGAGAAG  
 GTAAAATCTCTTAAAAAATTGAAAAAAGGAGCCACTATTGCAATTCC  
 AAATGATGCAACAAATGGTAGCCGTGCATTGTATGTCCTTCAGTCAG  
 CAGGTTTAAATCAAATTGAATGTTTCTGGTAAGAAGGTTGCAACAGTT  
 GCTAATATCACATCTAATAAAAAGGATATTAATATTCAGGAGTTAGA  
 TGCGAGTCAAACACCACGTGCACTCAAAGATGTAGATGCAGCTATTA  
 TTAATAATACATACATTGAGCAAGCTAATTTAAAACCTTCAGATGCT  
 ATCTTTGTTGAGAAATCAGATAAAAATTCAAAACAATGGATTAATAT  
 CATTGCGGGACGTAAAAAATTGGAAAAAGCAAAGAACGCTAAAGCT  
 ATCCAAGCTATCTTGGATGCTTATCACACAGATGAAGTGAAAAAAGT  
 TATCAAAGATACTTCAGCTGATATTCCACAATGGTAA

MILITSYGIISLSQKLREFIMKLKHIVLGLALTLLGVTFNSNQEVSASSTSS  
 KVVKVGVMTFSDTEKARWDKIEKLVGDKAKIKFTEFTDYTQPNQATAN  
 KDVDINAFQHYNFLENWNKENKKNLIPLEKTYLAPIRIYSEKVKSLKKL  
 KKGATIAIPNDATNGSRALYVLQSAGLIKLNVSGKKVATVANITSNKKDI  
 NIQELDASQTPRALKDVDAAIINNTYIEQANLKPDAIFVEKSDKNSKQW  
 INIAGRKNWKKQKNAKAIQAILDAYHTDEVKKVIKDT SADIPQW\*

ID-74

Clone 3-6

ATGTCAAATCAATATGATTATATCGTTATTGGTGGAGGTAGTGCAGG  
 CAGTGGTACCGCTAATAGGGCAGCCATGTATGGAGCAAAGTCCTGT  
 TAATTGAAGGTGGACAAGTAGGTGGAACCTTGTGTTAACTTAGGTTGT  
 GTACCTAAGAAAATCATGTGGTATGGTGCACAAGTTTCTGAGACACT  
 CCATAAGTATAGTTCAGGTTATGGTTTTGAAGCCAATAATCTTAGTTT  
 TGATTTTACTACTCTAAAAGCTAATCGCGATGCTTACGTGCAGCGGTC  
 TAGACAGTCGTATGCCGCTAATTTTGAAGCGTAATGGGGTTCGAAAAGA

---

FIG. 1 CONT'D

TTGATGGATTGCTCGTTTTATTGATAACCATACTATTGAAGTGAATG  
 GTCAGCAATATAAAGCTCCTCACATTACTATTGCAACAGGTGGACAC  
 CCTCTTTACCCTGATATTATTGGAAGTGAACCTGGTGAGACTTCTGAT  
 GATTTTTTTGGATGGGAGACCTTACCAAATTCTATATTGATTGTTGGG  
 GCGGGCTATATCGCGGCAGAACTTGCTGGAGTGGTTAATGAATTAGG  
 CGTTGAAACCCATCTTGCATTTAGAAAAGACCATATTCTACGCGGAT  
 TTGATGACATGGTAACAAGTGAGGTTATGGCTGAAATGGAGAAATCA  
 GGTATCTCTTTACATGCTAACCATGTACCTAAATCTCTTAAACGCGAT  
 GAAGGTGGCAAGTTGATTTTTGAAGCTGAAAATGGGAAAACGCTTGT  
 CGTTGATCGTGTAATATGGGCTATCGGCCGTGGACCAAATGTAGACA  
 TGGGACTTGAAAATACCGATATTGTTTTAAATGATAAAGATTATATC  
 AAAACAGATGAATTTGAGAATACTTCTGTAGATGGCGTGTATGCTAT  
 TGGAGATGTTAATGGGAAAATTGCCTTGACACCGGTAGCAATTGCAG  
 CAGGTCGTCGCTTATCAGAAAGACTTTTTAATCATAAAGATAACGAA  
 AAATTAGATTACCATAATGTACCTTCAGTTATTTTTACTCACCTGTA  
 ATTGGGACGGTAGGACTTTCAGAAGCAGCAGCTATCGAGCAATTTGG  
 AAAAGATAATATCAAAGTCTATACATCAACTTTTACCTCTATGTATAC  
 GGCTGTTACCAGTAATCGCCAAGCAGTTAAGATGAAGCTCATAACCC  
 TAGGAAAAGAGGAAAAAGTTATTGGGCTTCATGGTGTGGTTATGGT  
 ATTGATGAAATGATTCAAGTTTTTCAGTTGCTATCAAATGGGGGC  
 TACTAAAGCAGACTTTGATGATACTGTTGCTATTCACCCAACCTGGATC  
 TGAGGAATTTGTTACAATGCGCTAA

MSNQYDYIVIGGGSAGSGTANRAAMYGAKVLLIEGGQVGGTCVNLGC  
 VPKKIMWYGAQVSETLHKYSSGYGFEANNLSFDFTTLKANRDAYVQRS  
 RQSYAANFERNGVEKIDGFARFIDNHTIEVNGQYKAPHITIAATGGHPLY  
 PDIIGSELGETSDDFFGWETLPNSILIVGAGYIAAELAGVVNELGVETHLA  
 FRKDHILRGFDDMVTSEVMAEMEKSGISLHANHVPKSLKRDEGGKLIFE  
 AENGKTLVVDRVIWAIGRGNVDMGLENTDIVLNDKDYIKTDEFENTS  
 DGVYAIGDVNGKIALTPVAIAAGRRLSERLFNHKDNEKLDYHNVPSVIF  
 THPVIGTVGLSEAAAIEQFGKDNKVVYTSFTSMYTAVTSNRQAVKMKLI  
 TLGKEEKVIGLHGVGYGIDEMIQGFVSAIKMGATKADFDDTVAIHPTGS  
 EEFVTMR\*

ID-75

Clone 3-51

ATGAGTATCAAAAAAAGTGTGATTGGTTTTTGCCTCGAAGCTGCAGC  
 ATTATCAATGTTTGCTTGTGTAGACAGTAGTCAATCTGTTATGGCTGC

FIG. 1 CONT'D

CGAGAAGGATAAAGTCGAAATTACGTGGTGGGCTTTTCCAACCTTTA  
 CTCAAGAAAAGGCTAAGGATGGAGTAGGTA CTTATGAGAAAAAAGT  
 CATCAAGGCTTTTGAAAAGAAAAATCCTAATATAAAAAGTAAAACTAG  
 AGACAATTGATTTACATCTGGACCTGAAAAAATCACTACAGCAATT  
 GAAGCAGGGACAGCACCTGATGTGCTTTTTGATGCACCAGGGCGAAT  
 TATTC AATATGGTAAAAATGGTAAATTAGCAGATTTGAATGATTTATT  
 TACAGACCAATTTATTAAGGATGTCAATAATAAGAACATCATTCAAG  
 CTTCTAAGTCTGGCGATAAAGCCTACATGTATCCAATAAGTTCTGCC  
 CTTTTATATGGCGTTCAATAAAAAAATGCTTAAAGATGCAGGAGTT  
 TTGAAACTTGTAAGAAGGTTGGACTACTAGTGATTTTGAAAAAGT  
 ACTAAAAGCACTAAAAATAAAGGCTATACACCAGGTTCAATTCTTTG  
 CAAACGGGCAAGGAGGAGATCAAGGACCACGTGCATTTTTTGCTAAT  
 CTTTATAGTGCTCCAATAACAGATAAAGAAGTAACAAAATATACCAC  
 TGACACTAAAAATTCTGTAAAATCAATGAAAAAATAGTTGAATGGA  
 TTAAGAAAGGCTACTTGATGAATGGGTCTCAGTATGATGGCTCAGCT  
 GACATTCAAA ACTTCGCCAATGGACAAACTGCTTTC ACTATCCTATG  
 GGCTCCAGCTCAACCAAAAACTCAAGCAAAATTATTAGAGTCAAGTA  
 AAGTGGATTACCTTGAAGTGCCATTCCCATCAGAAGATGGAAAACCA  
 GATTTAGAATACCTTGTTAATGGTTTTGCGGTCTTTAATAATAAAGAT  
 GAAAACAAAGTAAAAGCCTCTAAGAAATTTATCACTTTTATTGCTGA  
 TGATAAAAAATGGGGACCAAAAGATGTTATACGTACAGGTGCTTTCC  
 CAGTTAGAACATCATTTGGGGATCTTTATAAAGGTGATAAACGTATG  
 ATGAAGATTTCAAAATGGACTCAATATTATTACCATATTACAACAC  
 TATCGATGGATTTTCTGAAATGAGAACCTTATGGTTCCCAATGGTTCA  
 ATCTGTATCCAATGGTGATGAAAAACCAGCAGATGCTTTGAAAGACT  
 TACTCAAAAAGCAAATGATACCATTA AAAAAGCAGCTAAATAA

MSIKKSVIGFCLEAAALSMFACVDSSQSVMAAEKDKVEITWWAFPTFTQ  
 EKAKDGVGTYEKKVIKAFEKKNPNIKVKLETIDFTSGPEKITTAIEAGTAP  
 DVLFDAPGRIIQYGKNGKLADLNDLFTDQFIKDVNNKNIIQASKSGDKA  
 YMPISSAPFYMAFNKKMLKDAGVLKLVKEGWTTSDFEKVLKALKNK  
 GYTPGSFFANGQGGDQGPRAFFANLYSAPITDKEVTKYTTDTKNSVKSM  
 KKIVEWIKKGYLMNGS QYDGSADIQNFANGQTAFTILWAPAQPKTQAK  
 LLESSKVDYLEVPPFSEDGKPDLEYLVNGFAVFNNK DENKVKASKKFIT  
 FIADDKKWGPKDVIRTGAFPVRTSFGDLYKGDKRMMKISKWTQYYSY  
 YNTIDGFSEMRTLWFP MVQSVSNGDEKPADALKDFTQKANDTIKKA  
 \*

ID-76 (Same as ID-39)

Clone 3-56

ATGAGGAAACGTTTTTCCTTGCTAAATTTTATTGTTGTTACTTTTATTT  
 TCTTTTTCTTTATTCTTTTTCCGCTTTTTAAGGCCAAAGATTGTCAGGT

FIG. 1 CONT'D

TGTTTATGCAAGTTTTCAAGGAGATCATTGGGACATTTGTAACGCATT  
TGATTTTCCGTATTTACATCGCTTTGATCTCATTAAAGGTAAAGAAAA  
TCAACTTTACTTTATAGGTTGTACAATTGCTAACAGTAAAGCCTACAC  
TGAGGATTGGAGTGATAAAGGCCGAATTTTTGTTGCTCGTTTTAATAC  
TCAAACCATACATTGGAAGGATTGCAACAATTGCCTCAAACCTTTAT  
TAAAAAATCATGGATACTATGCCATTCAGGATGAAGGATATTCATTG  
ATTACTTCAGTAGAAGGGGTACTCAAACCTCACTTATCCAGAATTTTCT  
ACTACAGGCGACTGGCAATTAGAACGGCTTTTCGATGAGGAGACAAG  
CGATGTGGTGAAAGTGGATATTAATCAGGATGGTAAGGATGAGTATG  
TGATCATCCAAGGTTTTCATGGAGATCGTTTACGTATCTTCACTGAAG  
ATTTCCGGTCGAGAATTATTCATTATCCTGAAAAAACCCATTTGGTC  
ACGCTATTTGGAGTGGTCGTTTACTTAATCAGACTTGTTTCGTATTTCG  
GGTGCGATCAGAAAAAGCAGAATTAAGGCTTTTTCACTTTGTAGAT  
GGCACTTGGTTTCAGAATTAGTAGATGCAAAGCAGCTTCTAGTAA  
TGTCTTAGCTTTTGAAAAAGATGGAAAAGCTTATCTTTCTCAGCCAA  
TAACGGACGTGGCGAAGTTGCTCTTTATCAATTAGTAAAATAA

MRKRFSLLNFIVVTFIFFFIFLPLFKAKDCQVVYASFQGDHWDICNAFDF  
PYLHRFDLIKGENQLYFIGCTIANSKAYTEDWSDKGRIFVARFNTQNHT  
LEGLQLPQTLLKNHGYAIQDEGYSLITSVEGVLLKLTYPFSTTGDWQ  
LERLFDEETSDVVKVDINQDGKDEYVIIQGFHGDRLRIFTEDFGRELFHY  
PEKTPFGHAIWSGRLLNQTCFVFGWRSEKAELRLFHFVDGHLVSELVDA  
KAASSNVLAFEKDGKAYLFSANNGRGEVALYQLVK\*

FIG. 1 CONT'D

nucS1

Bgl II Eco RV

5'-cgagatctgatatctcacaaacagataacggcgtaaataag -3'

nucS2

Bgl II Sma I

5'-gaagatcttccccgggatcacaaacagataacggcgtaaataag -3'

nucS3

Bgl II Eco RV

5'-cgagatctgatatccatcacaaacagataacggcgtaaataag -3'

nucR

Bam HI

5'-cgggatccttatggacctgaatcagcgttgctc -3'

NucSeq

5'-ggatgctttgtttcaggtgtatc -3'

pTREP<sub>F</sub>5'-catgatatcgggtacctcaagctcatatcattgtccggcaatgggtgtgggctttttttagcggataa  
caatttcacac -3'pTREP<sub>R</sub>5'-gcggatccccgggcttaattaatgtttaaacactagtcgaagatctcgcgaattctcctgtgtgaaatt  
gttatccgcta -3'pUC<sub>F</sub>

5'-cgccaggggttttcccagtcacgac -3'

V<sub>R</sub>

5'-tcaggggggocggagcctatg -3'

V<sub>1</sub>

5'-tcgtatggtgtgtggaattgtg -3'

V<sub>2</sub>

5'-tccggctcgtatggtgtgtggaattg -3'

FIG. 2

pTREP-Nuc vectors allow cloning of genomic DNA into each frame with respect to the nuclease gene

(i)

pTREP1-nuc1 (EcoRV)	AAGTATCAGATCT-- <u>GATATC</u> --TCACAAACAGATAACGGCGTAAAT	Frame=+1
	..... ▲ .....	
pTREP1-nuc2 (Sma I)	AAGTATCAGATCTT <u>CCCCGGGA</u> -TCACAAACAGATAACGGCGTAAAT	Frame=+2
	..... ▲ .....	
pTREP1-nuc3 (EcoRV)	AAGTATCAGATCT-- <u>GATATCCATC</u> ACAAACAGATAACGGCGTAAAT	Frame=+3
	..... ▲ .....	
Nuclease Gene		TCACAAACAGATAACGGCGTAAAT

Cloning site is indicated by an arrow

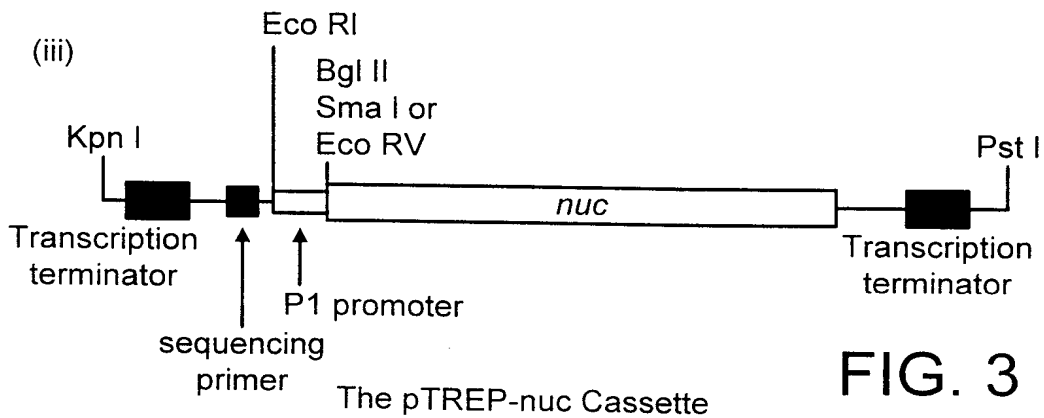
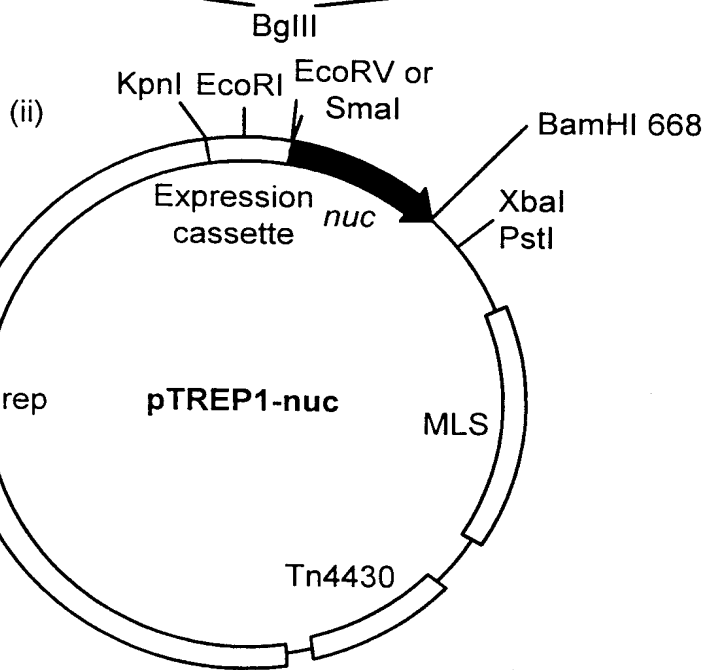


FIG. 3



GBS Vaccination - Trial 1

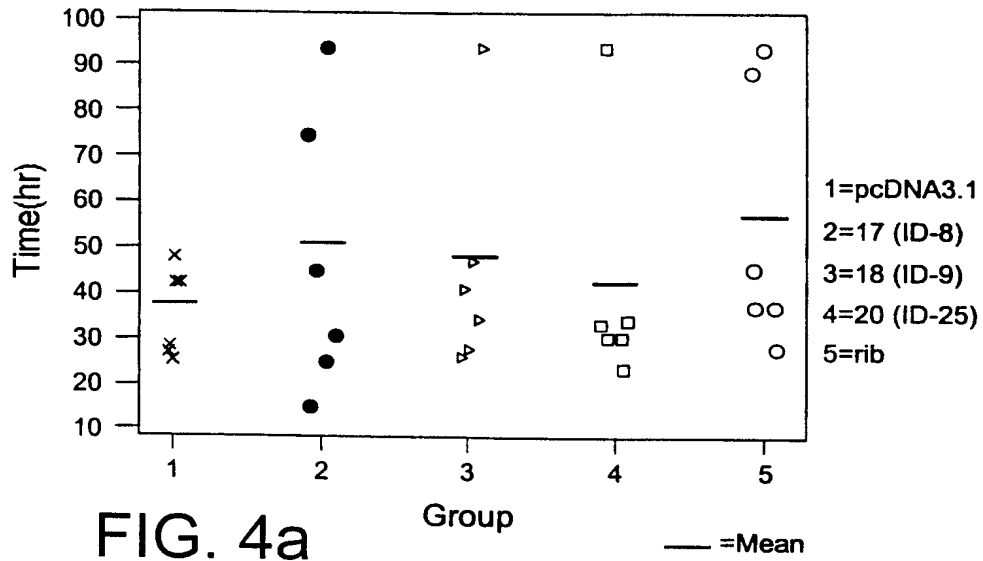


FIG. 4a

GBS Vaccination - Trial 2

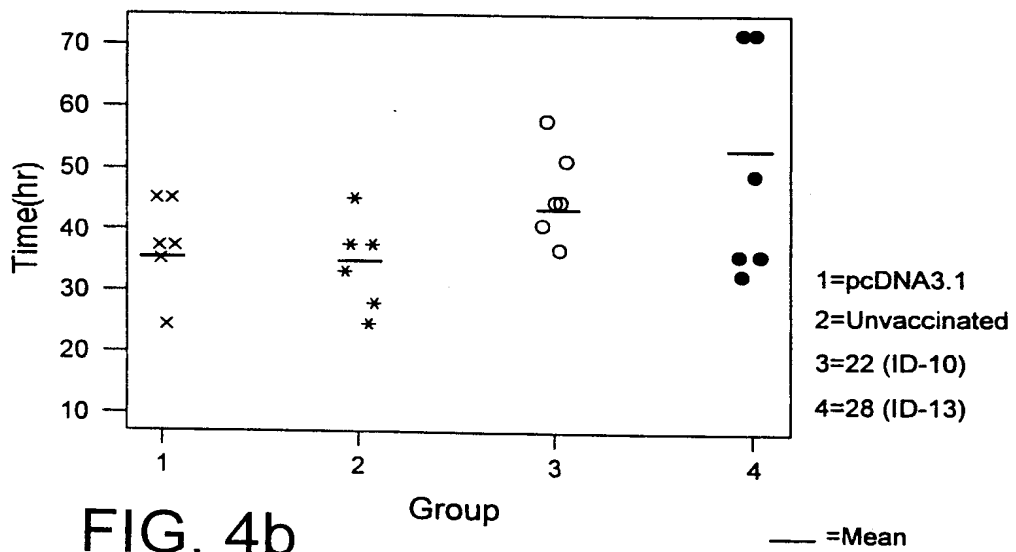


FIG. 4b

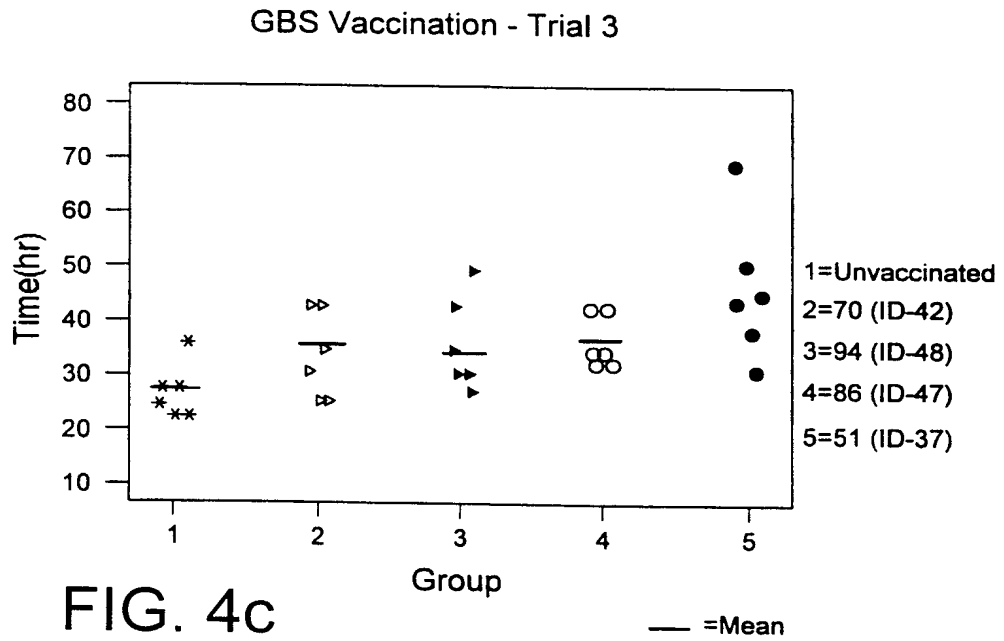


FIG. 4c

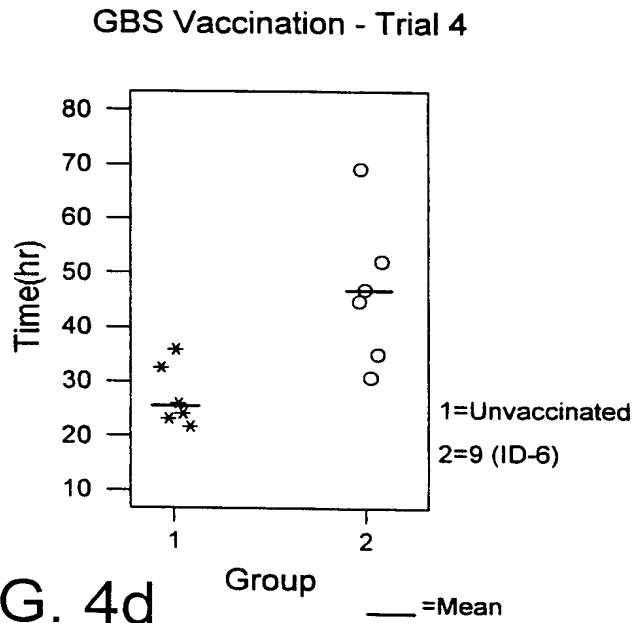


FIG. 4d

GBS Vaccination - Trial 6

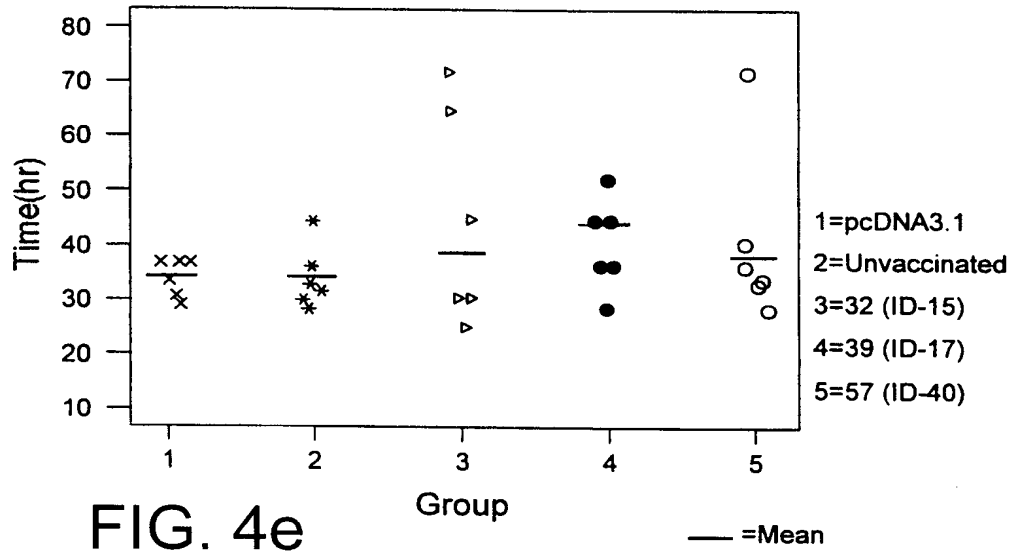


FIG. 4e

GBS Vaccination - Trial 2

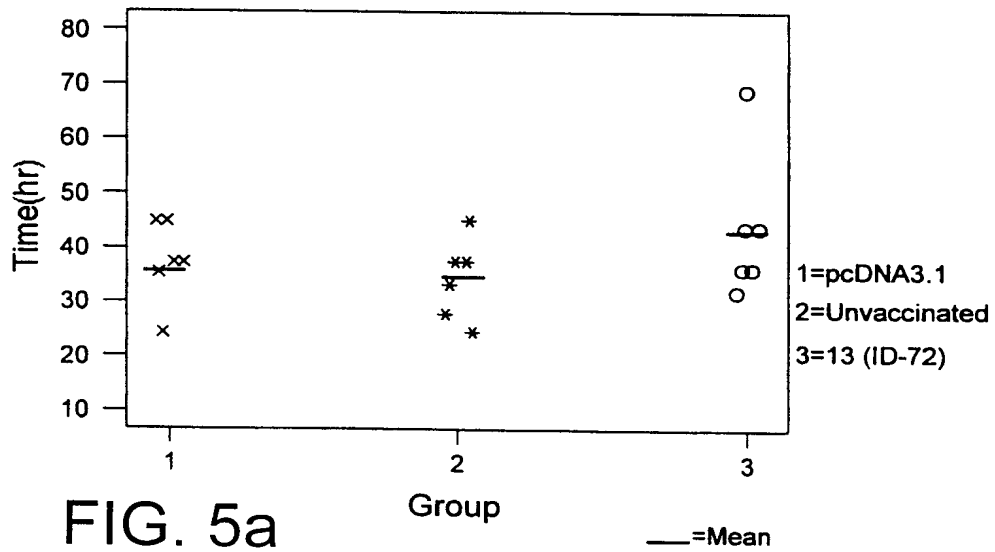


FIG. 5a

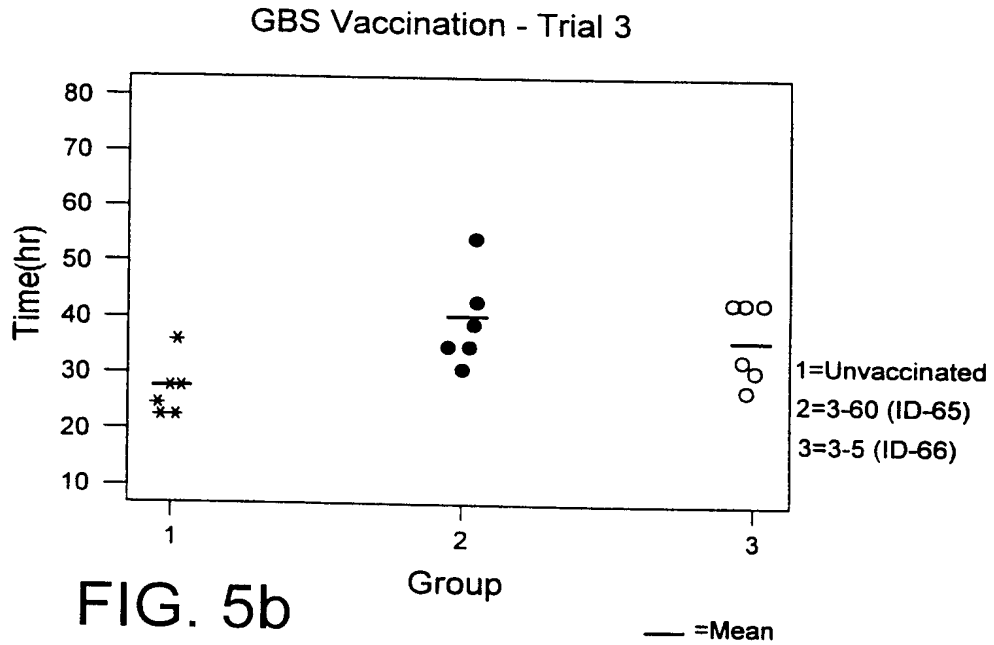


FIG. 5b

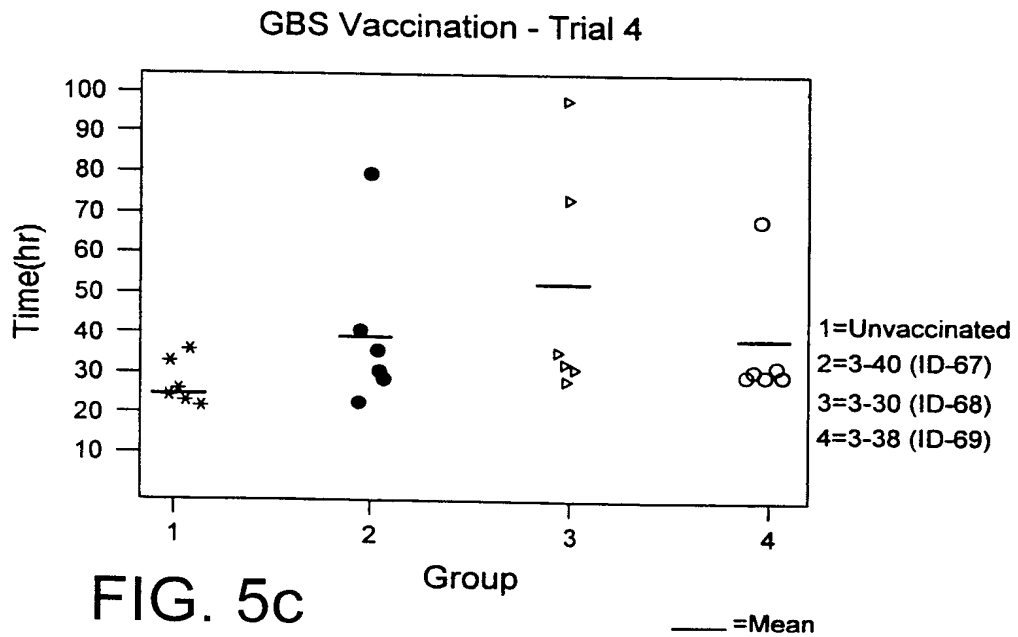


FIG. 5c

GBS Vaccination - Trial 5

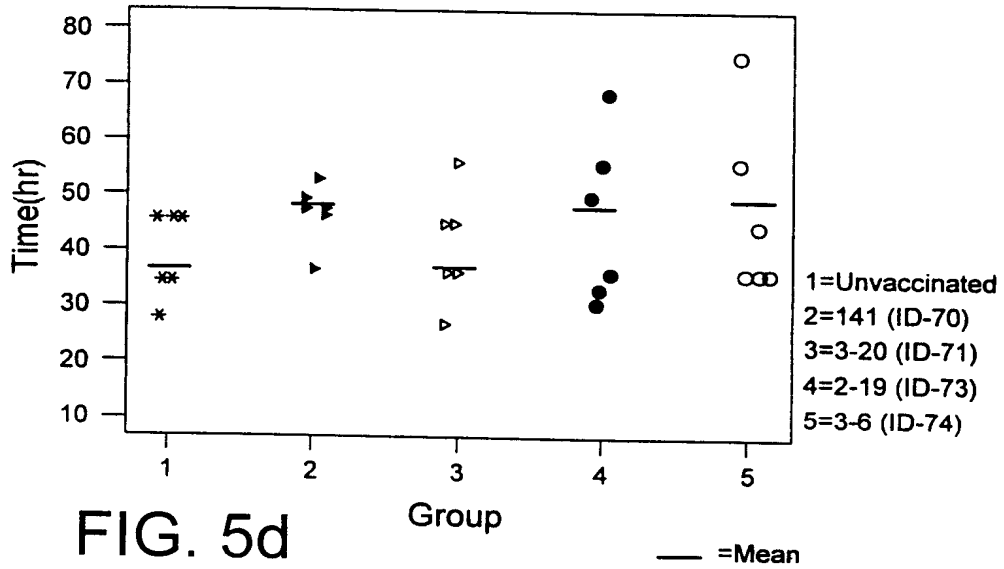


FIG. 5d

GBS Vaccination - Trial 6

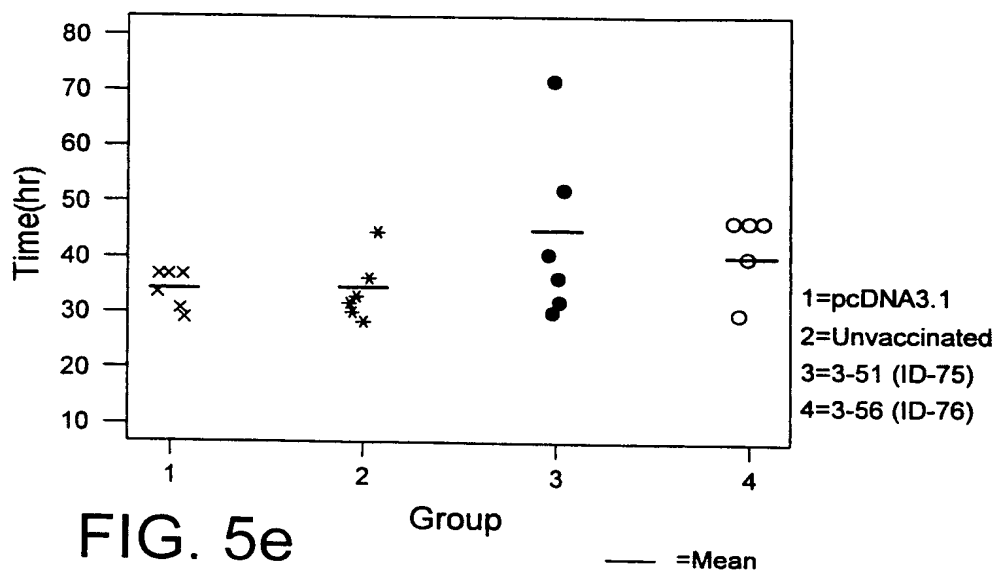


FIG. 5e

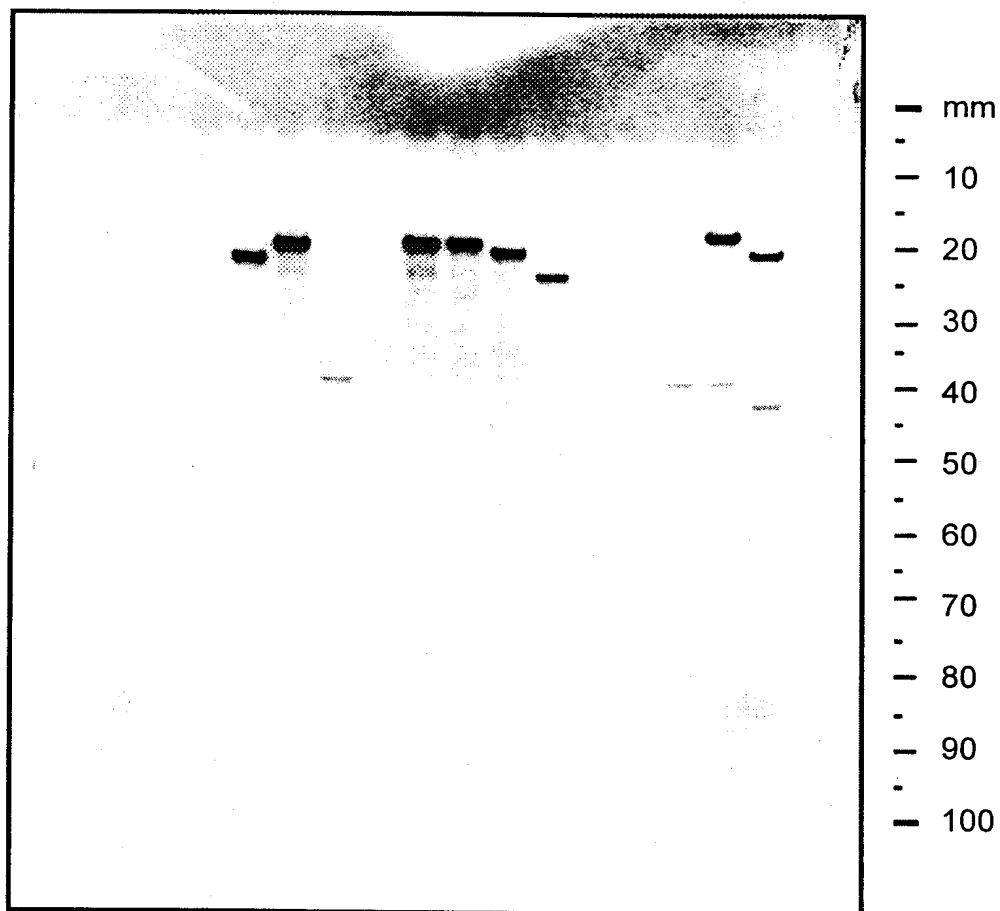


FIG. 6

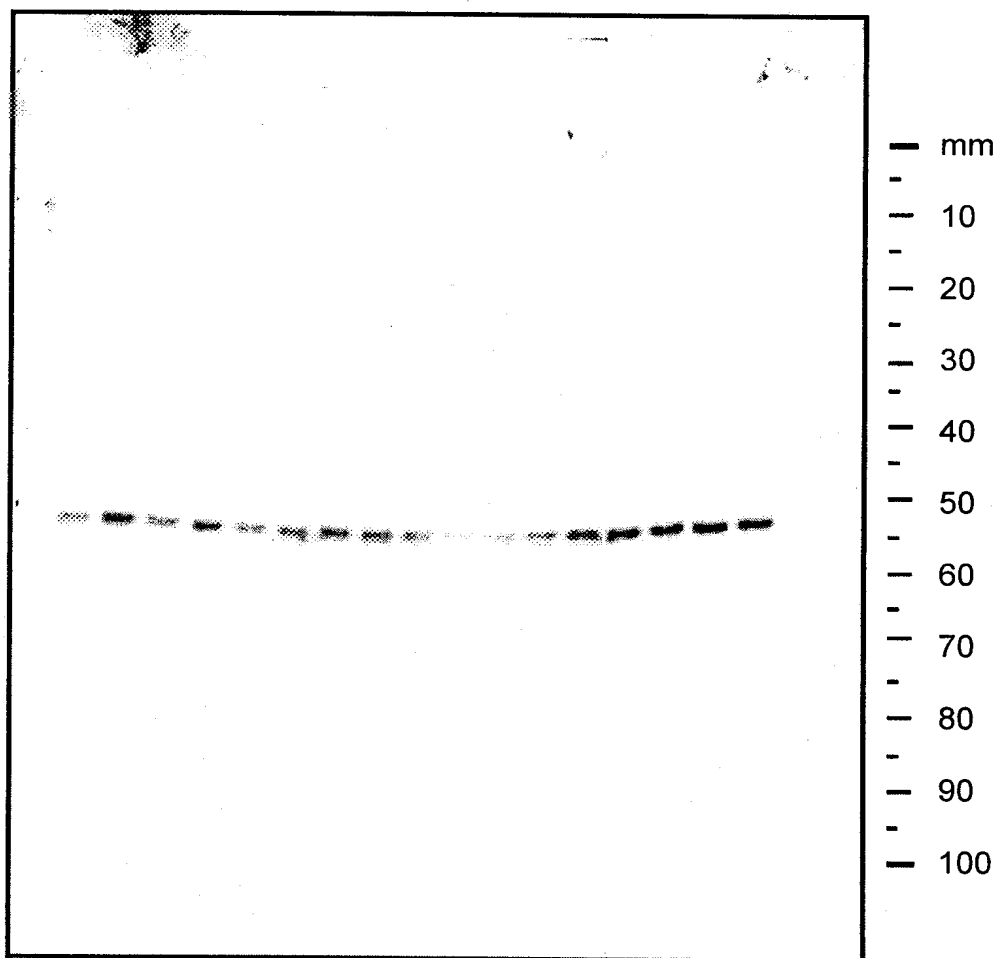


FIG. 7

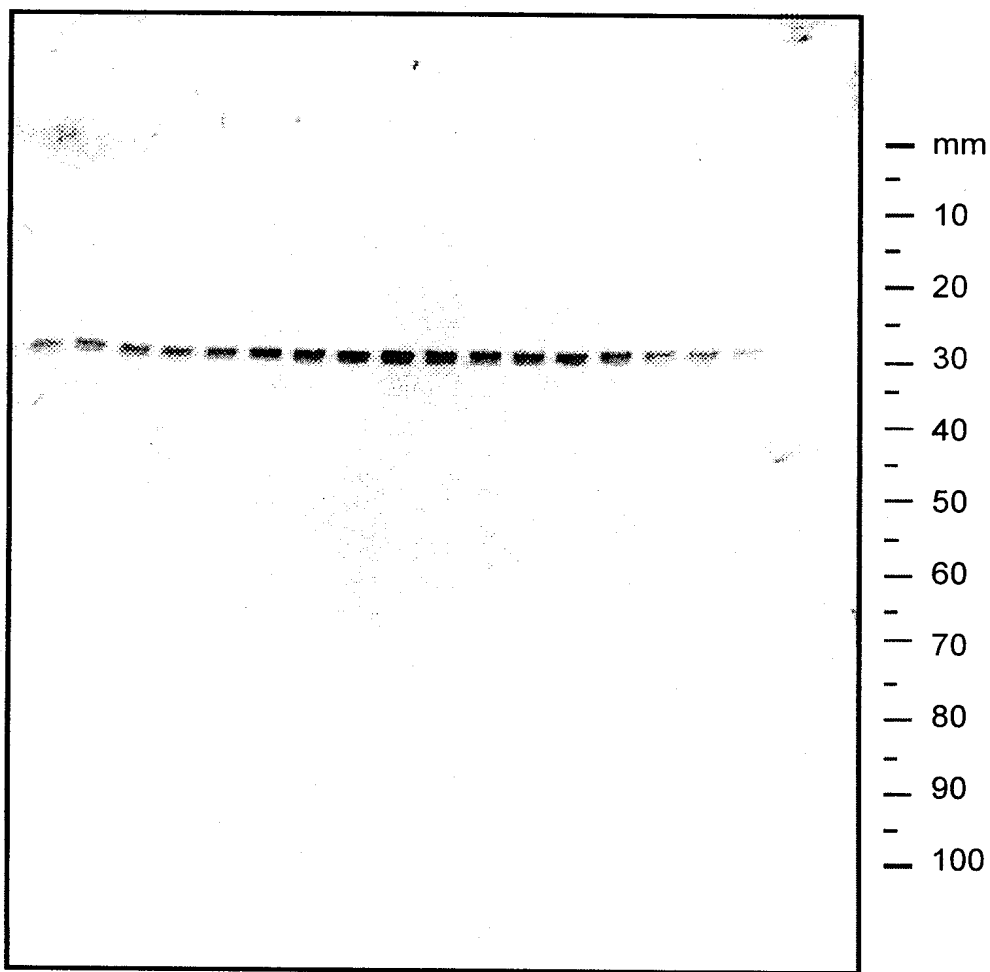


FIG. 8



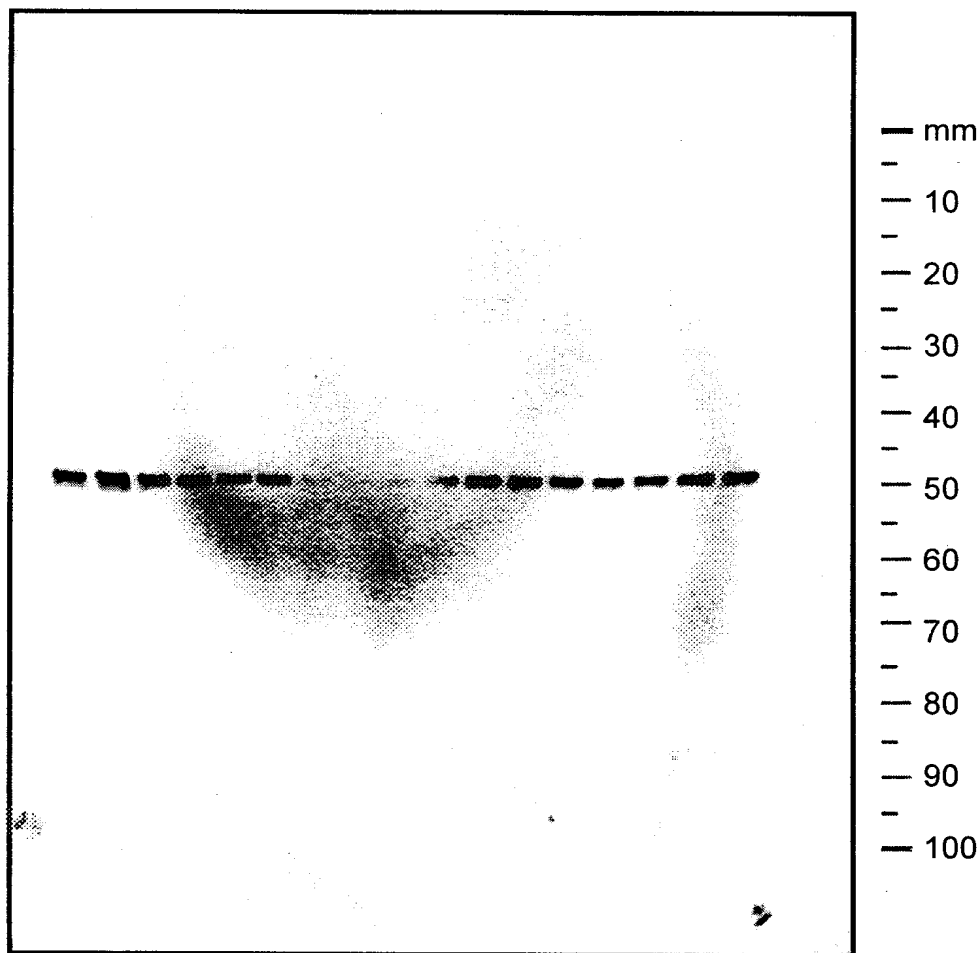


FIG. 9

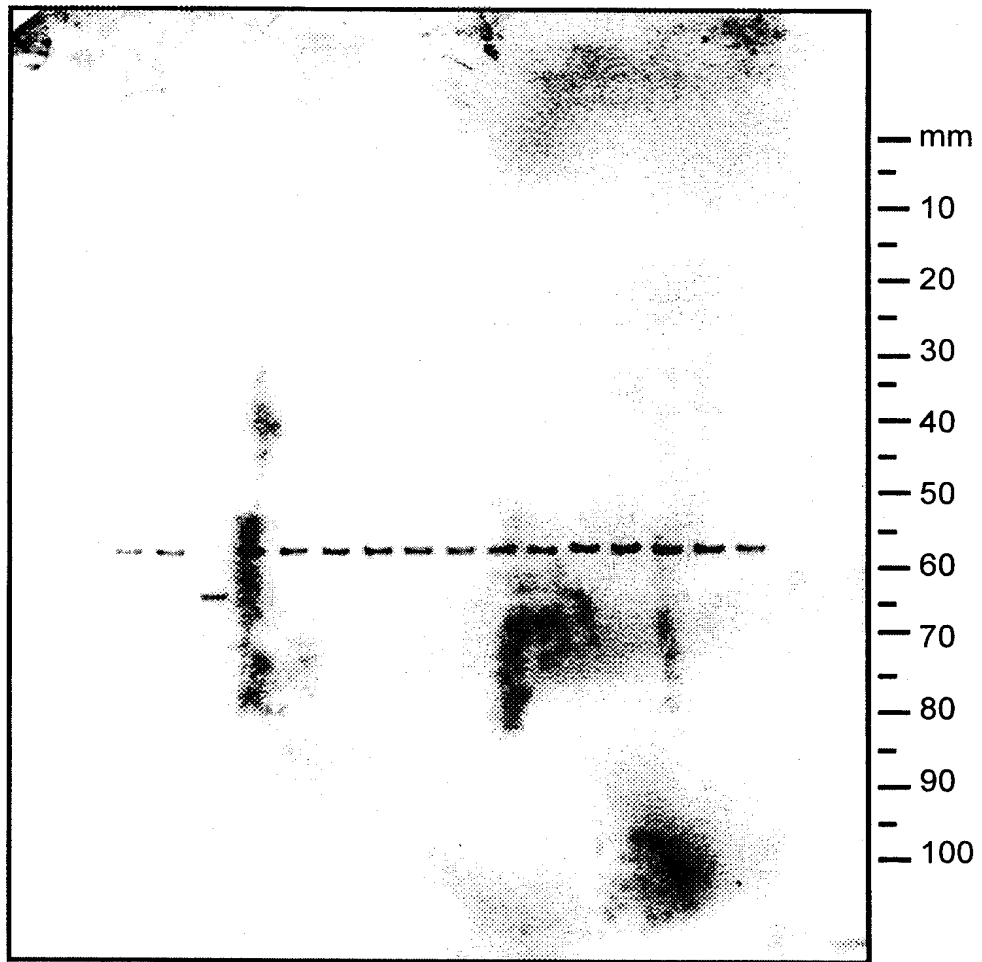


FIG. 10

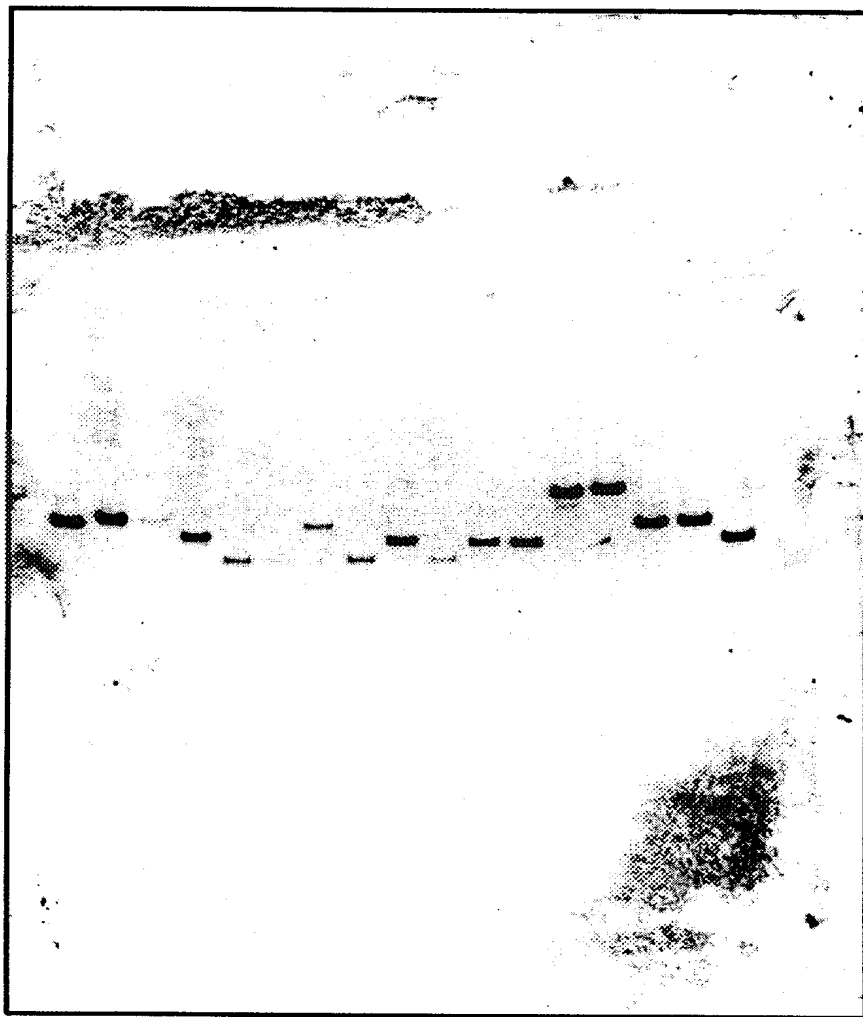


FIG. 11



## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

<p>(51) International Patent Classification <sup>7</sup> : C12N 15/31, 15/74, 15/62, 15/10, 9/16, 1/19, 1/21, C07K 14/315, 16/12, A61K 31/70, 39/09, G01N 33/53, 33/68, C12Q 1/68</p>	A3	<p>(11) International Publication Number: <b>WO 00/06736</b></p> <p>(43) International Publication Date: 10 February 2000 (10.02.00)</p>					
<p>(21) International Application Number: PCT/GB99/02444</p> <p>(22) International Filing Date: 27 July 1999 (27.07.99)</p> <p>(30) Priority Data:</p> <table border="0"> <tr> <td>9816335.5</td> <td>27 July 1998 (27.07.98)</td> <td>GB</td> </tr> <tr> <td>60/125,163</td> <td>19 March 1999 (19.03.99)</td> <td>US</td> </tr> </table> <p>(71) Applicant (for all designated States except US): MICROBIAL TECHNICS LIMITED [GB/GB]; 20 Trumpington Street, Cambridge CB2 1QA (GB).</p> <p>(72) Inventors; and (75) Inventors/Applicants (for US only): LE PAGE, Richard, William, Falla [GB/GB]; University of Cambridge, Dept. of Pathology, Tennis Court Road, Cambridge CB2 1QP (GB). WELLS, Jeremy, Mark [GB/GB]; Institute of Food Re- search, Norwich Laboratory, Norwich Research Park, Col- ney, Norwich NR4 7UA (GB). HANNIFFY, Sean, Bosco [IE/GB]; University of Cambridge, Dept. of Pathology, Ten- nis Court Road, Cambridge CB2 1QP (GB).</p> <p>(74) Agents: CHAPMAN, Paul, William et al.; Kilburn &amp; Strode, 20 Red Lion Street, London WC1R 4PJ (GB).</p>	9816335.5	27 July 1998 (27.07.98)	GB	60/125,163	19 March 1999 (19.03.99)	US	<p>(81) Designated States: CA, CN, JP, US, European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE).</p> <p><b>Published</b> <i>With international search report.</i></p> <p>(88) Date of publication of the international search report: 22 June 2000 (22.06.00)</p>
9816335.5	27 July 1998 (27.07.98)	GB					
60/125,163	19 March 1999 (19.03.99)	US					
<p>(54) Title: NUCLEIC ACIDS AND PROTEINS FROM GROUP B STREPTOCOCCUS</p>							
<p>(57) Abstract</p> <p>Novel protein antigens from Group B <i>Streptococcus</i> are described, together with nucleic acid sequences encoding them. Their use in vaccines and screening methods is also described.</p>							

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# INTERNATIONAL SEARCH REPORT

International Application No <b>PCT/GB 99/02444</b>
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<b>A. CLASSIFICATION OF SUBJECT MATTER</b>					
IPC 7	C12N15/31	C12N15/74	C12N15/62	C12N15/10	C12N9/16
	C12N1/19	C12N1/21	C07K14/315	C07K16/12	A61K31/70
	A61K39/09	G01N33/53	G01N33/68	C12Q1/68	

According to International Patent Classification (IPC) or to both national classification and IPC

**B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)

IPC 7 C12N C07K A61K G01N C12Q

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	DATABASE TREMBL E.M.B.L. Databases Accession Number: Q54914, 1 November 1996 (1996-11-01) POBBIELSKI A ET AL: "ORF 1 AND ORF2 5' REGION" XP002133342 97.2% identity in 141 aa overlap with SeqIdNo.12 abstract  <div style="text-align: center;">                         ---                          -/--                     </div>	3, 4

Further documents are listed in the continuation of box C.

Patent family members are listed in annex.

° Special categories of cited documents :

- \*A\* document defining the general state of the art which is not considered to be of particular relevance
- \*E\* earlier document but published on or after the international filing date
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- \*O\* document referring to an oral disclosure, use, exhibition or other means
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- \*X\* document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
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Date of the actual completion of the international search

17 March 2000

Date of mailing of the international search report

↑ 1. 04. 00

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International Application No  
PCT/GB 99/02444

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	<p>WO 98 18930 A (HUMAN GENOME SCIENCES INC ;CHOI GIL H (US); HROMOCKYJ ALEX (US); J) 7 May 1998 (1998-05-07) SP0020: 51.9% identity in 262 aa overlap with SeqIdNo.133 -&amp; DATABASE GENESEQ E.M.B.L. Databases Accession Number: W55078, 2 October 1998 (1998-10-02) CHOI G ET AL: "Streptococcus pneumoniae SP0020 protein" XP002133369 51.9% identity in 262 aa overlap with SeqIdNo.133 abstract</p>	3-18,23
P,X	<p>--- WO 99 16882 A (MEDIMMUNE INC) 8 April 1999 (1999-04-08) -&amp; DATABASE GENESEQ E.M.B.L. Databases Accession Number: Y05766, 8 April 1999 (1999-04-08) LUTTICKEN R ET AL : "Streptococcal adhesion mediator protein Lmb" XP002133343 99.7% identity in 306 aa overlap with SeqIdNo.12 abstract</p>	1-18,23
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A	<p>--- MICHEL J L ET AL: "Cloned alpha and beta C-protein antigens of group B Streptococci elicit protective immunity" INFECTION AND IMMUNITY,US,AMERICAN SOCIETY FOR MICROBIOLOGY. WASHINGTON, vol. 59, no. 6, June 1991 (1991-06), page 2023-2028-2028 XP002107260 ISSN: 0019-9567 the whole document</p>	1-18,23
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## INTERNATIONAL SEARCH REPORT

national Application No  
PCT/GB 99/02444

## C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
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## INTERNATIONAL SEARCH REPORT

national Application No  
PCT/GB 99/02444

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	<p>DATABASE TREMBL E.M.B.L. databases Accession Number: P94374, 1 May 1997 (1997-05-01) YOSHIDA K ET AL: "HOMOLOGOUS TO MANY ATP-BINDING TRANSPORT PROTEINS" XP002133346 30.2% identity in 235 aa overlap with SeqIdNo.82 abstract</p> <p style="text-align: center;">---</p>	
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T	<p>WO 99 42588 A (BIOCHEM VACCINS INC ;BRODEUR BERNARD R (CA); CHARLEBOIS ISABELLE ()) 26 August 1999 (1999-08-26)</p> <p style="text-align: center;">-----</p>	

# INTERNATIONAL SEARCH REPORT

International application No.  
PCT/GB 99/02444

## Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)

This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1.  Claims Nos.:  
because they relate to subject matter not required to be searched by this Authority, namely:
  
2.  Claims Nos.:  
because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
  
3.  Claims Nos.:  
because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

## Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

see additional sheet

1.  As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
  
2.  As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
  
3.  As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:  
  
1-18 and 23 (all partially) as relating to inventions 1, 6, 10, 13, 35, 41, 62, 63 and 67
  
4.  No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

### Remark on Protest

- The additional search fees were accompanied by the applicant's protest.
- No protest accompanied the payment of additional search fees.

## FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Invention 1: claims 1-18 and 23 (all partially)

A Group B Streptococcus protein having a sequence as depicted in SeqIdNo.2, a fragment, derivative or variant of said protein; a nucleic acid molecule comprising or consisting of SeqIdNo.1, a nucleic acid molecule complementary to said sequence, a nucleic acid molecule encoding for the a derivative or fragment of said protein; a vector comprising said nucleic acid molecule and afferent recombinant DNA practices; an antibody to said protein; an immunogenic composition comprising said protein or said nucleic acid and applications thereof; a method or kit of detection of Group B Streptococcus comprising said protein, said antibody, or said nucleic acid molecule; a method of determining whether said protein represents a potential antimicrobial target which comprises inactivating said protein and determining whether Group B Streptococcus is still viable.

2. Inventions 2-69: claims 1-18 and 23 (all partially)

Idem as subject 1 but limited to each of the polynucleotide and polypeptide sequences as depicted in SeqIdNo:3-137, wherein invention 2 is limited to SeqIdNo:3 and SeqIdNo:4, invention 3 is limited to SeqIdNo:5 and SeqIdNo:6, ..., invention 58 is limited to SeqIdNo:115, ..., and invention 69 is limited to SeqIdNo:136 and 137.

3. Inventions 70: claims 19-22 (all totally)

A method for screening for DNA encoding bacterial cell envelope associated or surface antigens in gram positive bacteria comprising a reporter vector including the nucleotide sequence encoding the mature form of the staphylococcus nuclease gene and an upstream promoter region with DNA from a gram positive bacterium; said method wherein the reporter vector is one of the pTREP1-nuc vectors; said method wherein the gram positive bacterium is Group B Streptococcus, Streptococcus pneumoniae, Staphylococcus aureus or pathogenic group A streptococci; said vector which is one of the pTREP1-nuc vectors

For the sake of conciseness, the first and 70th subject-matters are explicitly defined, the other subject-matters are defined by analogy to the subject-matter of invention 1.

## INTERNATIONAL SEARCH REPORT

Information on patent family members

International Application No

PCT/GB 99/02444

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